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Genome-wide association analysis identifies novel blood pressure loci and offers biological insights into cardiovascular risk

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Discovery and validation of 107 blood pressure loci from UK Biobank offers novel biological insights into cardiovascular risk

Short title: Novel blood pressure loci in UK Biobank

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Abstract:

Elevated blood pressure is the leading heritable risk factor for cardiovascular disease worldwide. We report genetic association of blood pressure (systolic, diastolic, pulse pressure) among UK Biobank participants of European ancestry with independent replication in other cohorts, leading to discovery and validation of 107 novel loci. We also identify new independent variants at 11 previously reported blood pressure loci. Combined with results from a range of *in-silico* functional analyses and wet bench experiments, our findings highlight new biological pathways for blood pressure regulation enriched for genes expressed in vascular tissues and identify potential therapeutic targets for hypertension. Results from genetic risk score models raise the possibility of a precision medicine approach through early lifestyle intervention to offset the impact of blood pressure raising variants on future cardiovascular disease risk.

Elevated blood pressure is a strong, heritable and modifiable driver of risk for stroke and coronary artery disease and a leading cause of global mortality and morbidity^{1,2}. In most populations blood pressure rises with age and by older ages over 50% of the population has hypertension^{3,4}. Raised blood pressure is heritable and arises from a complex interplay of lifestyle exposures and genetic background⁵⁻⁸. To date, studies including genome-wide meta-analyses of up to 2.5 million HapMap imputed variants across multiple studies, and analyses of bespoke or exome content, have identified 163 genetic variants of mostly modest or weak effect on blood pressure at 122 loci⁹⁻¹³. Here, we report association analyses between three blood pressure traits (systolic, diastolic and pulse pressure) and genetic variants among the first ~150,000 UK Biobank participants, with independent replication in large international consortia and other cohorts, providing new biological insights into blood pressure regulation.

UK Biobank is a prospective cohort study of 500,000 men and women aged 40-69 years with extensive baseline phenotypic measurements according to a standardized protocol (including blood pressure by a semi-automated device: Omron HEM-7015IT digital blood pressure monitor), stored biological samples (including DNA)¹⁴, and follow-up by electronic health record linkage¹⁵. Participants were genotyped using a customised array (including GWAS and exome content) and with genome-wide imputation based on 1000 Genomes and UK10K sequencing data^{16,17}.

Our study design is summarised in **Fig. 1**. Briefly, of the 152,249 UK Biobank participants with genotype data, after quality measures and exclusions (see Methods Online), we study 140,886 unrelated individuals of European ancestry with two seated clinic blood pressure measurements (**Supplementary Table 1**). We carry out genome-wide association study (GWAS) analyses of systolic, diastolic and pulse pressure using single-variant linear regression under an additive model, based on ~9.8 million single nucleotide variants (SNVs) with minor allele frequency (MAF) $\geq 1\%$ and imputation quality score (INFO) > 0.1 . We then consider for replication SNVs with $P < 1 \times 10^{-6}$ and take forward the sentinel SNV (i.e. with lowest P -value) at each locus, with a locus being defined by linkage disequilibrium (LD) $r^2 < 0.2$, within a 1Mb interval. We similarly analyse exome content for variants with MAF $\geq 0.01\%$, including rare variants, taking into replication the sentinel SNV ($P < 1 \times 10^{-5}$) from loci that are non-overlapping ($r^2 < 0.2$) with the GWAS findings. Overall we took the sentinel SNVs from 240 loci into replication ($r^2 < 0.2$ and $> 500\text{kb}$ from previously reported blood pressure SNVs and not annotated to previously reported blood pressure genes): 218 from GWAS and 22 from the exome analysis (GWAS variants from an additional 17 novel loci could not be taken into replication due to the absence of the variant or a proxy in the replication resources (**Supplementary Tables 2 and 3**).

The replication resources comprise a large BP meta-analysis consortium and further cohorts with 1000 Genomes data for the GWAS findings (**Supplementary Table 4**), and large blood pressure exome consortia meta-analyses, both with individuals of European ancestry. We use $P < 5 \times 10^{-8}$ to denote genome-wide significance in the combined (discovery and replication) meta-analyses, also requiring evidence of support ($P < 0.01$) in the replication data alone and concordant direction of effect. Additionally, we take forward for replication potential secondary signals at 51 previously reported blood pressure loci (excluding the HLA region). We note that the replication P -value threshold of $P < 0.01$ is more stringent than a range of

thresholds calculated according to False Discovery Rate (FDR) which gives FDR thresholds of $0.03 < P < 0.04$ (see Supplementary Methods).

To better understand the functional consequences of our new discoveries as well as previously reported variants, we carry out a series of *in silico* investigations including expression Quantitative Trait Locus (eQTL) analyses, tissue and DNASE hypersensitivity site enrichment and pathway analyses (**Supplementary Fig. 1**). We also test for long-range regulatory interactions (Hi-C) and investigate metabolomics signatures associated with our sentinel SNVs. Finally, we undertake experimental analysis of gene expression in relevant vascular tissue for selected putative functional SNVs.

RESULTS

Discovery and validation of genetic variants at novel loci

Of the 240 not previously reported loci taken forward to replication, we validate 107 novel loci at $P < 5 \times 10^{-8}$, of which 102 derive from the GWAS analysis replicated and meta-analysed in a total of 330,956 individuals (**Table 1a**; **Supplementary Fig. 2a-c**; **Supplementary Fig. 3a**), and a further five are from the exome analysis validated in a total of 422,604 individuals from the combined meta-analysis (**Table 1b** and **Supplementary Fig. 3b**; **Supplementary Tables 5 and 6**). Most SNVs also show association with hypertension in the UK Biobank data, for example 93 of the 107 validated novel sentinel SNVs are nominally significant ($P < 0.01$) (**Supplementary Table 7**).

Our results for systolic, diastolic and pulse pressure are shown in **Supplementary Figs. 2a,b,c** respectively. The most significant association signal for systolic pressure, which rises with age is with rs112184198 near *PAX2* ($P = 3.6 \times 10^{-18}$); for diastolic pressure, which plateaus in middle age, with rs76326501 near *METTL21A-AC016735.1* ($P = 3.6 \times 10^{-18}$); and rs3889199 near *FGGY* ($P = 1.8 \times 10^{-24}$) for pulse pressure, which increases with age and arterial stiffening¹⁸. However, as blood pressure traits are highly correlated, we unsurprisingly report considerable overlap in these findings (**Supplementary Fig. 4**). Many loci are associated with more than one blood pressure trait at genome-wide significance. For example, in the combined meta-analysis, 24 validated novel loci are associated with both systolic and diastolic pressure, 11 with both systolic and pulse pressure, one locus with both diastolic and pulse pressure and four loci (*NADK-CPSF3L*, *GTF2B*, *METTL21A-AC079767.3* and *PAX2*) are associated with all three traits (**Fig. 2**). We further note that many of the pulse pressure associated SNVs have opposing directions of effect for systolic and diastolic pressure, and are less likely to have strong associations with hypertension.

After conditional analysis on the sentinel SNV we identify five validated secondary SNVs in novel regions that are independently associated with blood pressure traits (**Table 2a**; **Supplementary Table 8**). We also note the existence of a rare validated potential secondary variant at the *NOX4* locus (rs56061986, MAF = 0.3%); although we do not claim this as an independent signal after conditioning on the sentinel variant, its relatively large effect on blood pressure remains (**Supplementary Table 8**). The contribution of our validated novel loci increases the percentage trait variance explained by ~1%, e.g. compared with 2.59% for previously reported SNVs alone, taken together, the validated novel and previously reported SNVs explain 3.56% of variance for systolic blood pressure, in an independent population.

For the first time in GWAS we report a signal at the angiotensin converting enzyme (*ACE*) locus ($P = 6.8 \times 10^{-14}$), from the renin-angiotensin system, a pathway which is targeted by current blood pressure treatments (ACE-inhibitors), as well as several other signals at known hypertension drug targets. These include *CACNA2D2* (rs743757, $P = 2.4 \times 10^{-10}$) targeted by calcium channel blockers, *MME* (rs143112823 in the RP11-439C8.2 locus, $P = 1.4 \times 10^{-14}$) targeted by omapatrilat for treating hypertension, *ADRA2B* (rs2579519 in the *GPAT2-FAHD2CP* locus, $P = 4.8 \times 10^{-12}$) targeted by beta blockers, *SLC14A2* (rs7236548, $P = 2.0 \times 10^{-18}$) targeted by the hypertension drug nifedipine, and phosphodiesterase 5A (*PDE5A*; rs66887589, $P = 3.4 \times 10^{-15}$) targeted by sildenafil for treating pulmonary hypertension.

Additionally, we evaluate our validated novel SNVs, where available, in cohorts of non-European ancestry^{12,13}, while recognising that these analyses are likely underpowered (**Supplementary Table 9**). For the GWAS SNVs, we find concordance in direction of effect ($P < 0.05$) for all three blood pressure traits for individuals of East Asian ancestry, and for diastolic pressure for South Asian ancestry. For the exome analyses, we find concordance in direction of effect among individuals of Hispanic ancestry. Despite small numbers, these findings point to cosmopolitan effects for many of the blood pressure associated variants.

A PhenoScanner¹⁹ search revealed that 27 of our 107 validated novel sentinel SNVs (or proxies; $r^2 \geq 0.8$) exhibit genome-wide significant associations (**Fig. 3a**) with other traits, including cardiovascular outcomes (e.g. coronary artery disease, myocardial infarction), cardiovascular risk factors (e.g. lipids, height, body mass index) and non-cardiovascular traits (e.g. lung function, cancer, Alzheimer's). Some of these associations may reflect genuine pleiotropic effects of variants. In some cases, such as for coronary artery disease, association with blood pressure may either be due to pleiotropy, or reflect the fact that elevated blood pressure lies on the causal pathway²⁰.

Associations at previously reported loci

In the conditional analyses, we identify 22 secondary SNVs (17 common, one rare and four low-frequency variants) that are conditionally independent of the blood pressure associated SNVs at 16 previously reported loci (**Table 2b; Supplementary Tables 10 and 11**). One rare variant (rs138582164, MAF=0.1%) in the *CDH17* locus anticipated to act as an exonic stop/gain mutation at the *GEM* gene is associated with a relatively large effect on pulse pressure (3.5 mm Hg per allele copy, **Table 2b**). At three previously reported loci (*EBF1*, *PDE3A*, *JAG1*) we identify multiple independent secondary SNVs in addition to the previously reported SNVs (**Supplementary Table 10**).

The UK Biobank data show support ($P < 0.01$) for 119 of the 122 previously reported blood pressure loci (159 of 163 SNVs) for one or more blood pressure traits (**Supplementary Fig. 2 a-c; Supplementary Table 12**). Thus we do not show support in UK Biobank for only four previously reported SNVs, one of which (rs11066280, *RPL6-ALDH1*) was identified from a GWAS of East Asian ancestry²¹ and may indicate ancestry-specific effects. We compare the MAF and effect sizes in UK Biobank with the published results of previously reported variants (**Supplementary Figure 5**), indicating consistency of results between the two sources of data.

We also examine findings for low-frequency and rare gene mutations previously reported to be associated with monogenic hypertension disorders²² and included on the UK Biobank gene array. Even within a large single study, there is still a lack of power for testing the impact of rare variants and it remains inconclusive as to whether any monogenic mutations also affect blood pressure levels within the general population. From the look-up results obtained within the UK Biobank data (**Supplementary Table 13**), there is suggestion that the variant with the lowest *P*-value (rs387907156; *KLH3*; MAF=0.02%) has a large effect on blood pressure (8.2 mm Hg per allele (SE=4.1); *P* = 0.046 and 5.6 mm Hg (SE=2.6); *P* = 0.048 for systolic and pulse pressure respectively).

Functional analyses

We annotate the 107 validated novel loci to 212 genes (based on LD $r^2 \geq 0.8$) and seek putative function from *in silico* analyses of our novel and previously reported loci, as well as undertaking gene expression experiments for selected SNVs in relevant vascular tissue. Candidate genes with the strongest supporting evidence are indicated in the last column of Table 1 with an indication of the supporting data source. All genome wide-significant variants in LD ($r^2 > 0.8$) for (a) validated novel loci and (b) previously reported loci, ranked by supporting evidence are annotated in **Supplementary Table 14**. Of the 107 validated novel sentinel SNVs only three are Indels, all other variants are single nucleotide polymorphisms (SNPs). We identify non-synonymous SNVs at 13 of the 107 validated novel loci, including three non-synonymous novel sentinel SNVs (rs1250259 at *FN1* locus, rs78648104 at *TFAP2D* and rs7127805 at *CRACR2B* locus) (**Supplementary Table 15**). Furthermore three of the 13 validated novel loci contain non-synonymous SNVs that are predicted to be damaging (ANNOVAR) in *TFAP2D* (rs78648104), *NOX4* (rs56061986, see above) and *CCDC141* (rs17362588, reported to be associated with heart rate²³) (**Fig. 3a**). Beyond the coding regions we identify 29 novel associated SNVs in 3'UTRs which are predicted to significantly weaken or cause loss of miRNA regulation by altering the recognition motif in seven genes, and strengthen or create target sites for miRNA binding in 13 genes (based on miRNASNP db, **Supplementary Table 15**).

Our expression Quantitative Trait locus (eQTL) analysis (based on GTEx data) shows that many novel loci contain variants with eQTLs across a range of different tissues (**Supplementary Table 16**). Of the 107 validated novel loci, 59 contain variants with eQTLs in at least one tissue. We observe arterial tissue as the tissue having the largest number of loci with eQTLs (**Supplementary Fig. 6**). Our follow-up targeted *in-silico* analysis reveals six novel loci with eQTLs in arterial tissue (**Supplementary Table 15**). For example, the GTEx tibial artery eQTL in *SF3A3* (rs4360494) shows strong *in silico* supporting evidence, including an arterial DNase I site within which the major C allele removes a predicted AP-2 binding site (**Supplementary Fig. 7**). Hence we prioritised this gene for *in vitro* functional analysis (see below).

By considering all loci together from both validated novel and previously reported loci, our analysis using DEPICT identifies enrichment of expression across 31 tissues and cells (**Supplementary Fig.8; Supplementary Table 17**), with greatest enrichment in the arteries (*P*

= 1.9×10^{-6} , false discovery rate (FDR) < 1%). We use FORGE to investigate and identify significant (FDR, $P < 0.05$) cell type specific enrichment within DNase I hypersensitive sites in a range of tissues including dermal and lung microvascular endothelial cell types, and cardiac fibroblasts (**Supplementary Fig. 9**). For a set of curated candidate regulatory SNVs from validated novel loci (see Supplementary Methods), widespread enrichment is found in microvascular endothelium, aortic smooth muscle, aortic fibroblasts, vascular epithelium, heart and skin (**Supplementary Fig. 9**). In addition, we identify significant enrichment of histone marks in a wide range of cell types, including strong enrichment seen for H3K4Me3 (an activating modification found near promoters) marks in umbilical vein endothelial cells (HUVEC) (**Supplementary Fig. 10**). To explore expression at the level of cardiovascular cell types specifically, we use Fantom5 reference transcript expression data (see Methods Online) to cluster the 212 genes annotated to our 107 validated novel loci according to tissue specificity (**Supplementary Fig. 11**), with the significantly clustered genes forming four tissue-specific clusters, including a vascular smooth muscle cell (VSMC) and fibroblast cluster, an endothelial cell cluster (including probable endothelial cells in highly vascularised tissues), and a combined vascular cell cluster.

Additionally, Ingenuity pathway analysis and upstream transcriptional analysis show enrichment of canonical pathways implicated in cardiovascular disease, including those targeted by antihypertensive drugs, such as the alpha-adrenergic, CXCR4, endothelin signalling and angiotensin receptor pathways (**Supplementary Table 18**). In keeping with vascular mediation of genetic influence we identify diphenyleneiodonium, an inhibitor of flavin-containing oxidases, including NAD(P)H oxidase, which is reported to reverse endothelial dysfunction (and hypertension) in a rat model²⁴.

In order to identify long range target genes of non-coding variants, we use chromatin interaction (Hi-C) data from HUVEC, as enhancers and silencers often form chromatin loops with their target promoter. In most loci the strongest promoter interaction involves a gene in high LD with the SNV but for 21 loci we find a distal potential target gene (**Supplementary Table 15**). Ingenuity pathway analysis of the distal genes shows the greatest enrichment in regulators of cardiac hypertrophy.

We further evaluate pleiotropy using the Genomic Regions Enrichment of Annotations Tool (GREAT) to study enrichment of mouse phenotype and human disease ontology terms across all our validated novel and previously reported loci. These highlight cardiovascular system abnormalities and vascular disease as the most highly enriched terms (**Fig. 3b & 3c**).

Collectively evidence from eQTLs, DEPICT, DNase I sites, histone marks, Hi-C data and ontological analyses indicates predominant vascular and cardiovascular tissue involvement for genes within the blood pressure associated loci. For example, aggregating all loci together in the DEPICT analysis, we observe greatest enrichment in arterial tissue, which has the largest proportion of novel loci having variants with eQTLs.

We also look for association of our validated sentinel SNVs with metabolomic signatures. Three novel SNVs within the *NOX4*, *KCNH4* and *LHFPL2* loci show significant associations (family-wise error rate < 5%) with lipoprotein sub-fractions from ¹H Nuclear Magnetic

Resonance (NMR) spectroscopy analysis of 2,000 Airwave study samples (**Supplementary Tables 19 and 20**). The results for these variants suggest a link between blood pressure regulation and lipid metabolism. Eleven SNVs (including at *LHFPL2* locus) show association (family wise error rate < 5%) with metabolites in blood or urine from the publicly available “Metabolomics GWAS Server” resource based on mass spectrometry^{25,26} (**Supplementary Table 20**), including sugar acids, sphingolipids, fatty acids, glycerophospholipids, organic acids and benzene derivatives.

Several genes and variants with putative function are highlighted in our *in silico* analysis as having biological support (e.g. eQTLs or nsSNVs) and those with novelty and tractability to laboratory investigation (e.g. expression in available tissue models) are prioritized. Sentinel variants in three genes are selected for experimental testing and successfully genotyped, each for at least 100 samples. We select *ADAMTS7* due to strong biological support (e.g. mouse knockout phenotype), *SF3A3* due to eQTLs and *NOX4* as it contains a rare nsSNV in addition to common variant associations. All three SNVs reached highly significant levels of association with blood pressure in the combined meta-analysis (**Table 1**): rs62012628 at *ADAMTS7* for diastolic pressure (0.238 mmHg per allele ± 0.03 , $P=5.1 \times 10^{-12}$, $N=244,143$); rs4360494 at *SF3A3* for pulse pressure (0.278 mmHg ± 0.03 , $P=3.7 \times 10^{-16}$, $N=307,682$); rs2289125 at *NOX4* for pulse pressure (-0.377 mmHg ± 0.04 , $P=9.1 \times 10^{-22}$, $N=282,851$). We use quantitative polymerase chain reaction (qPCR) to study the impact of these sentinel variants on gene expression in human vascular smooth muscle (VSMCs) and endothelial cells (ECs) (see Methods Online). For *SF3A3*, the major C allele of sentinel variant rs4360494 associated with increased pulse pressure is also associated with *SF3A3* expression in human VSMCs, although this SNV is not related to expression in endothelial cells (**Supplementary Fig. 12a**); and the T allele of SNV rs62012628 in *ADAMTS7*, associated with lower diastolic pressure, is associated with reduced *ADAMTS7* expression in human VSMCs (**Supplementary Fig. 12b**). Moreover, we find that the minor A allele of sentinel SNV rs2289125 at the *NOX4* locus correlates with increased *NOX4* expression in ECs though not VSMCs (**Supplementary Fig. 12c**). Our study thus finds evidence for novel cis-eQTLs in *ADAMTS7* and *NOX4* in addition to validating the previously reported GTEx eQTL in *SF3A3*, and supports the vascular expression of these genes.

Genetic risk of increased blood pressure, hypertension and cardiovascular outcomes

We create an unbiased genetic risk score (GRS) (**Supplementary Table 21**) to evaluate, in an independent cohort (Airwave, see Methods Online), the impact of the combination of our validated novel and previously reported loci on blood pressure levels and risk of hypertension. When compared with the lowest quintile of the distribution of the GRS, individuals >50 years in the highest quintile have sex-adjusted mean systolic pressure higher by 9.3 mm Hg (95% CI 6.9 to 11.7 mm Hg, $P=1.0 \times 10^{-13}$) and an over two-fold higher risk of hypertension (OR 2.32 95% CI 1.76 to 3.06; $P=2.8 \times 10^{-9}$) compared with individuals in the lowest quintile of the GRS distribution (**Fig. 4; Supplementary Table 22**). Similar results were obtained from GRS associations with blood pressure and hypertension within UK Biobank (**Supplementary Table 23**). In UK Biobank – based on self-reported health data, record linkage to Hospital Episode Statistics and mortality follow-up data (**Supplementary Table 24**) – we show that the GRS is associated with increased risk of stroke, coronary heart disease and all cardiovascular outcomes, comparing the upper and lower fifths of the GRS distribution, with sex-adjusted

odds ratios of 1.34 (95% CI 1.20 to 1.49, $P = 1.5 \times 10^{-7}$), 1.38 (95% CI 1.30 to 1.47, $P = 4.3 \times 10^{-23}$) and 1.35 (95% CI 1.27 to 1.42, $P = 1.3 \times 10^{-25}$) respectively (**Fig. 4; Supplementary Table 25**). Results are also provided for incident-only cases (**Supplementary Table 26**).

DISCUSSION

A key attribute of this study is the combination of a large, single discovery sample with standardized blood pressure measurement and a dense 1000 Genomes imputation strategy (UK 10K enhanced 1000G imputation), yielding a high quality dataset of ~9.8 million variants for study¹⁶. This is the largest genetic association analysis for blood pressure to date taking advantage of major international consortia for parallel replication of common and low-frequency variants, based in total on data from 330,956 individuals and exonic SNVs in a total of 422,604 individuals²⁷. This strategy resulted in the discovery of 107 robustly validated novel loci for blood pressure traits. In previous large-scale blood pressure genome-wide association scans we estimated that an effective doubling of sample size from a discovery cohort of 70,000 to 140,000 individuals with ~2.5 million imputed variants would double the number of validated loci, resulting in an estimated ~30 additional loci for blood pressure traits²⁷. Here we find over three times that number, taking advantage of UK Biobank's standardized approach to data collection, biobanking, genotyping and enhanced imputation strategy. Despite its size, our study is still under-powered to find low-frequency variants and the vast majority of our findings are common variants, with similarly modest or small effect sizes as previously reported validated variants (**Supplementary Fig. 13**). Our GWAS, which was restricted to $MAF \geq 1\%$, only identified four novel sentinel SNVs of low-frequency ($1\% \leq MAF < 5\%$) and our Exome analysis, despite allowing for rare variant discovery, did not identify any rare novel sentinel SNVs. The only rare and low-frequency variants identified were secondary SNVs within previously reported loci. The lack of rare variant discovery could also be due to the challenge of detecting rare variants from imputed data, in contrast to the recent Exome-chip studies which identified some novel rare SNVs from genotyped data^{11,12}. There may be greater potential for identifying rare variants from the future release of genetic data for all 500,000 UK Biobank participants.

Our findings point to new biology as well as highlighting novel gene regions in systems that have previously been implicated in the genetics of blood pressure. Several of our validated novel loci affect atherosclerosis or vascular remodelling (*ADAMTS7*, *THBS2*, *CFDP1*) and exhibit locus pleiotropy in prior genome-wide association studies for coronary artery disease or carotid intimal-media thickness²⁸⁻³⁰ (**Fig. 3a** and **Fig. 5**). In previous work we have shown that expression of *ADAMTS7* is upregulated and increases vascular smooth muscle cell migration in response to vascular injury in relation to a distinct coronary artery variant (rs3825807 which is not in strong LD with our sentinel SNV; $r^2 = 0.17$)³¹. In endothelial cells *ADAMTS7* acts as a metalloproteinase to cleave thrombospondin-1 encoded by *THBS2* which leads to reduced endothelial cell migration and plays a role in neo-intimal repair in the vessel wall³¹. Our functional work indicates that the allele associated with lower diastolic pressure is also associated with lower *ADAMTS7* expression in human vascular smooth muscle cells; this fits with the murine knockout that exhibits reduced atherosclerosis. *SF3A3* is a splicing

factor with no prior links to blood pressure other than our reported association and eQTL. NOX4 has an established role in the endothelium where it enhances vasodilatation and reduces blood pressure in vivo³². At the CFDP1 locus our sentinel SNV is in high LD ($r^2 = 0.95$) with a variant previously associated with carotid intimal-medial thickness. Collectively our findings highlight a potential common mechanism among these genes in vascular remodelling that has previously been observed in small resistance arteries in essential hypertension³³.

We identify both common and rare variant associations at the novel NADPH oxidase 4 (*NOX4*) locus. This oxidase generates reactive oxygen species in the endothelium and may contribute to salt sensitive hypertension in the kidney and the vasculature³⁴⁻³⁶. We found that the allele of the common variant at *NOX4* locus correlates with increased tissue specific *NOX4* expression in endothelial cells rather than vascular smooth muscle cells (**Supplementary Figure 12c**). *NOX4* mediates endothelial cell apoptosis and facilitates vascular collagen synthesis contributing to endothelial dysfunction and arterial stiffness, and may explain the association with pulse pressure^{37,38}.

We identify several loci containing genes involved in vascular signalling and second messenger systems such as *PDE5A* and *PDE10A*³⁹⁻⁴¹. The phosphodiesterase *PDE5A* hydrolyses cyclic GMP and is inhibited by sildenafil which leads to vasodilatation⁴². This finding fits with our previous discoveries of a role for gene loci encoding elements of natriuretic peptide-nitric oxide pathway and guanylate cyclase signalling systems in blood pressure regulation^{21,43,44}. Our findings strengthen the case for evaluating the opportunity to repurpose *PDE5A* inhibitors for use in hypertension.

The importance of microvascular function is emphasised by the solute carrier transporters such as *SLC14A2* encoding a urea transporter, which has previously been linked to autosomal dominant Streeten type orthostatic hypotensive disorder⁴⁵ and blood pressure response to nifedipine, a calcium channel blocker antihypertensive drug⁴⁶. *SLC8A1* encodes a sodium calcium exchanger expressed in cardiomyocytes which alters cardiac contractility and hypertrophy and shows abnormal blood pressure in *SLC8A1* transgenic mice⁴⁷. Variants at *SLC35F1* have been previously associated with resting heart rate and ventricular dimensions which could contribute to blood pressure elevation⁴⁸.

We also identify loci that are involved in cardiovascular development (*GATA2*, *KIAA1462*, *FBN2*, *FN1* and *HAND2*) such as fibrillin 2 (*FBN2*) which overlaps in action with fibrillin 1 in development of the aortic matrix⁴⁹⁻⁵³. In addition, fibronectin expression is increased in hypertension and in atherosclerosis but it may also play a role in the development of the heart⁵³⁻⁵⁵.

Our analysis validates loci containing genes with prior physiological connection to blood pressure such as *BDNF*, *FAM208A*, and *CACNA2D2*⁵⁶⁻⁵⁸. The neurotrophin Brain Derived Neurotrophic Factor modulates angiotensin 11 in the brain to elevate blood pressure in experimental models and higher serum levels correlate with reduced risk of cardiovascular disease and mortality⁵⁶. In experimental models *FAM208A*, which is thought to be a transcription factor, is a strong candidate for a quantitative trait locus for blood pressure⁵⁸. The gene *CACNA2D2* encodes a subunit of the L-type calcium channel that is most abundantly

expressed in the atrium and in neurones and may be a target for negatively chronotropic and inotropic calcium channel antagonists which reduce blood pressure⁵⁹.

This is the first time long range genomic interactions have been sought using Hi-C for blood pressure, where the promoter region has a strong chromatin interaction with a novel SNV. One such gene is *EPAS1*, which is ~200kb away from the SNV (rs11690961). It encodes hypoxia-inducible factor 2alpha, which affects catecholamine homeostasis, protects against heart failure and mutations in the gene are associated with pulmonary hypertension⁶⁰. Another gene is *INHBA*, 1.3Mb away from the SNV (rs12531683), which is elevated in pulmonary hypertension and contributes to vascular remodelling by inducing expression of endothelin-1 and plasminogen activator inhibitor-1 in pulmonary smooth muscle cells⁶¹.

Our observation that the blood pressure genetic risk score is associated with 9-10 mm Hg higher blood pressure at age 50+ years when comparing the top vs bottom fifths of the distribution in an independent population has potential clinical and public health implications. The results are particularly striking when stratified by age, due to a significant interaction of the GRS with age (P ranging between 9.96×10^{-11} and 1.16×10^{-3} for interaction with continuous blood pressure traits, and $P = 0.012$ for hypertension). Measuring the genetic risk score in early life raises the possibility of adopting an early precision medicine approach to risk management through lifestyle intervention (i.e. reduced sodium intake, increased potassium intake, maintenance of optimal weight, low adult alcohol consumption and regular exercise)⁶²⁻⁶⁴. Indeed, studies of non-pharmacologic approaches to blood pressure control indicate that we might achieve 10 mm Hg or more reduction in systolic blood pressure through lifestyle measures alone⁶⁵. At the same time, recent evidence suggests that favorable lifestyle may offset the cardiovascular sequelae associated with high genetic risk⁶⁶. However, as the above data are observational, it is not certain to what extent adherence to lifestyle recommendations amongst high genetic risk individuals could result in favorable outcomes. Given the substantial effect of genetic risk score on blood pressure by middle-age, the potential for adopting early lifestyle intervention amongst individuals at high genetic risk, along with population-wide measures to lower blood pressure, warrants further study.

Since the completion of our study, another blood pressure GWAS has been recently published⁶⁷. This used UK Biobank data within a larger single-stage combined meta-analysis, reporting a total of 316 loci, including 241 loci identified from the meta-analysis involving UK Biobank that were not tested for validation, as no replication resource was available. Our study reports 107 validated novel loci, of which 32 are detected and validated for the first time in our analysis of UK Biobank. In addition, 75 sentinel SNVs are in LD ($r^2 \geq 0.2$) with the recently reported loci and we are able to validate at least 53 of these for the first time in our study. Furthermore we note that 49 of the reported loci from this recent study⁶⁷ did not validate in our large independent replication resource.

We describe 107 validated novel loci for blood pressure offering new biology, identifying potential new therapeutic targets and raising the possibility of a precision medicine approach to modify risk of hypertension and cardiovascular outcomes. In total this brings the number of combined validated novel and previously reported loci for blood pressure traits to 229,

representing a major advance in our understanding of the genetic architecture of blood pressure.

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Conflicts/Disclosures

MJC is Chief Scientist for Genomics England, a UK government company.

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Central analysis: HRW, CPC, HG, MRB, MPSL, MR, IT, BM, IK, EE.

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Replication study contributor: [Lifelines] NV, PVDH, HS, AMS; [GS:SFHS] JM, CH, DP, SP; [EGCUT] TE, MA, RM, AM; [PREVEND] PVDH, NV, RTG, SJLB; [ASCOT] HRW, MJC, PBM, PS, NP, AS, DS, ST; [BRIGHT] HRW, MJC, PBM, MB, MF, JC; [Airwave] HG, EE, MPSL, IK, IT, PE.

All authors critically reviewed and approved the final version of the manuscript.

765 **Table 1: Association results for the sentinel variant from each validated novel locus from (a) UK Biobank GWAS discovery and (b) UK**
766 **Biobank exome discovery.** Results are shown for the primary blood pressure trait with most significant association from the combined meta-
767 analysis.

(a) UK Biobank GWAS																	
Sentinel SNV in the locus					UK Biobank discovery				Replication			Combined					
Locus	Chr	Pos	rsID	EA	EA	Beta	SE	P	Beta	SE	P	N	Beta	SE	P	Traits	Candidate genes
Systolic blood pressure																	
<i>NADK-CPSF3L</i>	1	1,685,921	rs139385870	D	0.50	-0.394	0.07	1.9x10 ⁻⁸	-0.310	0.07	1.0x10 ⁻⁵	281,890	-0.352	0.05	1.3x10 ⁻¹²	DBP,PP	GNB1 ^{df} , NADK ^{efg}
<i>CELA2A</i>	1	15,798,197	rs3820068	A	0.81	0.497	0.09	2.4x10 ⁻⁸	0.367	0.08	5.3x10 ⁻⁶	310,776	0.425	0.06	1.1x10 ⁻¹²	PP	AGMAT ^{df} , CELA2A ^{efg}
<i>GTF2B</i>	1	89,360,158	rs10922502	A	0.62	-0.475	0.07	4.7x10 ⁻¹¹	-0.307	0.06	2.0x10 ⁻⁶	323,666	-0.382	0.05	2.2x10 ⁻¹⁵	DBP,PP	KYAT3 ^{bf} , GTF2B ^{defg}
<i>FOSL2</i>	2	28,635,740	rs7562	T	0.52	0.365	0.07	2.2x10 ⁻⁷	0.182	0.06	3.7x10 ⁻³	319,942	0.263	0.05	1.9x10 ⁻⁸		FOSL2 ^{efg}
<i>PRKD3</i>	2	37,517,566	rs13420463	A	0.77	0.504	0.08	1.4x10 ⁻⁹	0.244	0.07	7.3x10 ⁻⁴	330,307	0.356	0.05	7.0x10 ⁻¹¹	DBP	PRKD3 ^{efg}
<i>METTL21A-AC079767.3</i>	2	208,526,140	rs55780018	T	0.54	-0.426	0.07	1.7x10 ⁻⁹	-0.360	0.07	5.1x10 ⁻⁸	304,567	-0.391	0.05	5.9x10 ⁻¹⁶	DBP,PP	METTL21A ^{efg}
<i>RYK</i>	3	134,000,025	rs9859176	T	0.40	0.419	0.07	6.4x10 ⁻⁹	0.248	0.06	9.6x10 ⁻⁵	322,428	0.322	0.05	1.3x10 ⁻¹¹	DBP	RYK ^{efg}
<i>NPNT</i>	4	106,911,742	rs13112725	C	0.76	0.418	0.08	3.1x10 ⁻⁷	0.450	0.08	9.4x10 ⁻⁹	306,370	0.435	0.06	1.5x10 ⁻¹⁴	DBP	NPNT ^{dfg}
<i>TMEM161B</i>	5	87,514,515	rs10059921	T	0.08	-0.644	0.13	5.9x10 ⁻⁷	-0.417	0.12	7.9x10 ⁻⁴	298,543	-0.526	0.09	4.0x10 ⁻⁹		TMEM161B ^g
<i>FBN2</i>	5	127,868,199	rs6595838	A	0.30	0.483	0.08	2.0x10 ⁻¹⁰	0.236	0.07	4.5x10 ⁻⁴	328,401	0.344	0.05	7.6x10 ⁻¹²	DBP	FBN2 ^{defg}
<i>CASC15</i>	6	22,130,601	rs6911827	T	0.45	0.433	0.07	8.2x10 ⁻¹⁰	0.190	0.06	2.1x10 ⁻³	326,471	0.296	0.05	2.0x10 ⁻¹⁰	DBP	CASC15 ^{efg}
<i>TFAP2D</i>	6	50,683,009	rs78648104	T	0.92	-0.664	0.13	1.2x10 ⁻⁷	-0.329	0.11	4.0x10 ⁻³	305,426	-0.481	0.08	1.3x10 ⁻⁸	DBP	TFAP2D ^{aefg}
<i>MKLN1</i>	7	131,059,056	rs13238550	A	0.40	0.486	0.07	9.4x10 ⁻¹²	0.212	0.06	7.1x10 ⁻⁴	325,647	0.331	0.05	1.9x10 ⁻¹²	DBP	PODXL ^{cf} , MKLN1 ^{efg}
<i>HIPK2</i>	7	139,463,264	rs1011018	A	0.20	-0.441	0.09	6.1x10 ⁻⁷	-0.244	0.08	1.6x10 ⁻³	325,110	-0.329	0.06	1.5x10 ⁻⁸		TBXAS1 ^{cdf} , HIPK2 ^{cdefg}
<i>ZFAT</i>	8	135,612,745	rs894344	A	0.60	-0.384	0.07	6.8x10 ⁻⁸	-0.163	0.06	8.2x10 ⁻³	329,834	-0.258	0.05	3.2x10 ⁻⁸		ZFAT ^{dfg}
<i>PAX2</i>	10	102,604,514	rs112184198	A	0.10	-0.826	0.12	7.8x10 ⁻¹³	-0.532	0.10	1.3x10 ⁻⁷	323,791	-0.659	0.08	3.6x10 ⁻¹⁸	DBP,PP	PAX2 ^{cefg}
<i>MCF2L</i>	13	113,636,156	rs9549328	T	0.23	0.440	0.08	1.5x10 ⁻⁷	0.218	0.08	3.9x10 ⁻³	313,787	0.318	0.06	1.5x10 ⁻⁸	PP	MCF2L ^{defg}
<i>FERMT2</i>	14	53,377,540	rs9888615	T	0.29	-0.427	0.08	3.5x10 ⁻⁸	-0.236	0.07	4.3x10 ⁻⁴	326,235	-0.318	0.05	3.5x10 ⁻¹⁰		FERMT2 ^{defg}
<i>PPP2R5E</i>	14	63,928,546	rs8016306	A	0.80	0.454	0.09	2.5x10 ⁻⁷	0.250	0.07	7.9x10 ⁻⁴	329,869	0.335	0.06	3.7x10 ⁻⁹	DBP	PPP2R5E ^{efg}
<i>ABHD17C</i>	15	81,013,037	rs35199222	A	0.45	0.353	0.07	5.7x10 ⁻⁷	0.298	0.06	1.7x10 ⁻⁶	323,407	0.322	0.05	5.2x10 ⁻¹²	DBP	ABHD17C ^{efg}
<i>CFDP1</i>	16	75,331,044	rs11643209	T	0.42	-0.481	0.07	1.8x10 ⁻¹¹	-0.222	0.06	6.3x10 ⁻⁴	309,242	-0.339	0.05	1.8x10 ⁻¹²	PP	CFDP1 ^{bfg} , BCAR1 ^{def}
<i>CRK</i>	17	1,333,598	rs12941318	T	0.49	-0.317	0.07	6.2x10 ⁻⁶	-0.226	0.07	6.9x10 ⁻⁴	299,739	-0.269	0.05	2.5x10 ⁻⁸	PP	CRK ^{cdg}

ACOX1	17	73,949,045	rs2467099	T	0.22	-0.423	0.08	4.5x10 ⁻⁷	-0.216	0.07	3.6x10 ⁻³	326,401	-0.307	0.06	3.3x10 ⁻⁸		ACOX1 ^{dfg} , FBF1 ^{acef}
Diastolic blood pressure																	
chr1mb25	1	25,030,470	rs6686889	T	0.25	0.231	0.05	3.7x10 ⁻⁷	0.143	0.04	9.1x10 ⁻⁴	322,575	0.185	0.03	3.6x10 ⁻⁹		RUNX3 ^{df} , CLIC4 ^{cdef} , SRRM1 ^g
DNM3	1	172,357,441	rs12405515	T	0.56	-0.219	0.04	4.1x10 ⁻⁸	-0.118	0.04	1.6x10 ⁻³	328,543	-0.165	0.03	1.4x10 ⁻⁹		DNM3 ^{defg}
GPATCH2	1	217,718,789	rs12408022	T	0.26	0.226	0.05	5.9x10 ⁻⁷	0.172	0.04	6.7x10 ⁻⁵	320,983	0.198	0.03	2.4x10 ⁻¹⁰		GPATCH2 ^g
CDC42BPA	1	227,252,626	rs10916082	A	0.73	-0.222	0.04	5.3x10 ⁻⁷	-0.135	0.04	1.5x10 ⁻³	327,636	-0.177	0.03	8.4x10 ⁻⁹		CDC42BPA ^{efg}
WNT3A	1	228,191,075	rs2760061	A	0.47	0.235	0.04	3.7x10 ⁻⁹	0.225	0.04	1.1x10 ⁻⁸	312,761	0.230	0.03	2.1x10 ⁻¹⁶	SBP	WNT9A ^{df} , WNT3A ^{efg}
SDCCAG8	1	243,471,192	rs953492	A	0.46	0.293	0.04	1.2x10 ⁻¹³	0.153	0.04	4.6x10 ⁻⁵	325,253	0.220	0.03	7.4x10 ⁻¹⁶		SDCCAG8 ^{bcefg}
ADCY3	2	25,139,596	rs55701159	T	0.89	0.382	0.06	1.1x10 ⁻⁹	0.193	0.06	1.6x10 ⁻³	321,052	0.285	0.04	7.2x10 ⁻¹¹	SBP	ADCY3 ^{efg}
SLC8A1	2	40,567,743	rs4952611	T	0.58	-0.200	0.04	8.0x10 ⁻⁷	-0.114	0.04	4.6x10 ⁻³	309,395	-0.157	0.03	4.0x10 ⁻⁸		SLC8A1 ^{cdefg}
AC016735.1	2	43,167,878	rs76326501	A	0.91	0.426	0.07	4.3x10 ⁻¹⁰	0.413	0.07	1.5x10 ⁻⁹	318,127	0.419	0.05	3.6x10 ⁻¹⁸	SBP	PRKCE ^{df} , HAAO ^g
GPAT2-FAHD2CP	2	96,675,166	rs2579519	T	0.63	-0.259	0.04	1.7x10 ⁻¹⁰	-0.137	0.04	6.7x10 ⁻⁴	311,557	-0.197	0.03	4.8x10 ⁻¹²		ADRA2B ^{cdf} , TCF7L1 ^{cef} , FAHD2CP ^{efg}
TEX41	2	145,646,072	rs1438896	T	0.30	0.288	0.04	2.1x10 ⁻¹¹	0.187	0.04	4.3x10 ⁻⁶	329,278	0.234	0.03	2.0x10 ⁻¹⁵	SBP	TEX41 ^g
CCDC141	2	179,786,068	rs79146658	T	0.91	-0.375	0.07	5.8x10 ⁻⁸	-0.245	0.07	4.2x10 ⁻⁴	321,318	-0.311	0.05	2.4x10 ⁻¹⁰		CCDC141 ^{afg}
TMEM194B	2	191,439,591	rs7592578	T	0.19	-0.271	0.05	8.9x10 ⁻⁸	-0.212	0.05	1.7x10 ⁻⁵	304,672	-0.240	0.04	9.5x10 ⁻¹²	SBP	NAB1 ^g
TNS1	2	218,668,732	rs1063281	T	0.60	-0.231	0.04	1.2x10 ⁻⁸	-0.172	0.04	1.4x10 ⁻⁵	315,354	-0.200	0.03	1.3x10 ⁻¹²	SBP	TNS1 ^{cefg}
CAMKV-ACTBP13	3	49,913,705	rs36022378	T	0.80	-0.265	0.05	6.3x10 ⁻⁸	-0.140	0.05	3.9x10 ⁻³	319,983	-0.202	0.03	4.7x10 ⁻⁹		CAMKV ^{efg}
CACNA2D2	3	50,476,378	rs743757	C	0.14	0.313	0.06	2.9x10 ⁻⁸	0.184	0.05	5.1x10 ⁻⁴	328,836	0.245	0.04	2.4x10 ⁻¹⁰		CACNA2D2 ^{dfg} , C3orf18 ^{def}
FAM208A	3	56,726,646	rs9827472	T	0.37	-0.207	0.04	3.6x10 ⁻⁷	-0.148	0.04	1.7x10 ⁻⁴	323,058	-0.177	0.03	4.3x10 ⁻¹⁰		FAM208A ^{efg}
RP11-439C8.2	3	154,707,967	rs143112823	A	0.09	-0.484	0.07	2.9x10 ⁻¹²	-0.295	0.08	2.3x10 ⁻⁴	297,343	-0.403	0.05	1.4x10 ⁻¹⁴	SBP	MME ^{defg}
SEN2	3	185,317,674	rs12374077	C	0.35	0.203	0.04	8.3x10 ⁻⁷	0.127	0.04	1.2x10 ⁻³	327,513	0.163	0.03	9.2x10 ⁻⁹		SEN2 ^{efg}
PDE5A	4	120,509,279	rs66887589	T	0.52	-0.296	0.04	5.7x10 ⁻¹⁴	-0.140	0.04	2.1x10 ⁻⁴	324,397	-0.215	0.03	3.4x10 ⁻¹⁵		FABP2 ^{cf} , PDE5A ^{defg}
POC5	5	75,038,431	rs10078021	T	0.63	-0.223	0.04	4.7x10 ⁻⁸	-0.105	0.04	9.2x10 ⁻³	314,172	-0.164	0.03	1.3x10 ⁻⁸		POC5 ^{efg}
CPEB4	5	173,377,636	rs72812846	A	0.28	-0.232	0.04	1.6x10 ⁻⁷	-0.186	0.04	2.4x10 ⁻⁵	312,601	-0.209	0.03	2.2x10 ⁻¹¹		C5orf47 ^{ef} , CPEB4 ^g
PKHD1	6	51,832,494	rs13205180	T	0.49	0.218	0.04	3.7x10 ⁻⁸	0.123	0.04	1.1x10 ⁻³	325,419	0.168	0.03	7.0x10 ⁻¹⁰		PKHD1 ^{cefg}
PDE10A	6	166,178,451	rs147212971	T	0.06	-0.421	0.08	2.3x10 ⁻⁷	-0.289	0.09	9.4x10 ⁻⁴	296,010	-0.360	0.06	1.6x10 ⁻⁹		PDE10A ^{defg}
SLC35F1	6	118,572,486	rs9372498	A	0.08	0.459	0.07	5.4x10 ⁻¹⁰	0.231	0.07	5.6x10 ⁻⁴	330,625	0.334	0.05	1.8x10 ⁻¹¹	SBP	SLC35F1 ^{dfg}
SNX31	8	101,676,675	rs2978098	A	0.54	0.212	0.04	6.9x10 ⁻⁸	0.122	0.04	1.4x10 ⁻³	324,424	0.165	0.03	1.5x10 ⁻⁹		SNX31 ^{efg}

RP11-273G15.2	8	144,060,955	rs62524579	A	0.53	-0.202	0.04	2.8x10 ⁻⁷	-0.140	0.05	2.2x10 ⁻³	268,645	-0.175	0.03	3.8x10 ⁻⁹		CYP11B1 ^{cdf} , CYP11B2 ^{cfg}
MTAP	9	21,801,530	rs4364717	A	0.55	-0.218	0.04	3.5x10 ⁻⁸	-0.136	0.04	2.9x10 ⁻⁴	327,173	-0.175	0.03	1.3x10 ⁻¹⁰		MTAP ^{acefg}
BDNF	11	27,728,102	rs11030119	A	0.31	-0.211	0.04	7.0x10 ⁻⁷	-0.119	0.04	3.3x10 ⁻³	330,002	-0.163	0.03	2.9x10 ⁻⁸		BDNF ^{cdefg}
MYEOV	11	69,079,707	rs67330701	T	0.09	-0.415	0.07	7.8x10 ⁻⁹	-0.314	0.08	3.8x10 ⁻⁵	276,760	-0.367	0.05	2.1x10 ⁻¹²	SBP	MYEOV ^{efg}
RP11-321F6.1	15	66,869,072	rs7178615	A	0.37	-0.207	0.04	3.8x10 ⁻⁷	-0.152	0.04	1.0x10 ⁻⁴	318,076	-0.179	0.03	2.6x10 ⁻¹⁰		LCTL ^{efg}
ADAMTS7	15	79,070,000	rs62012628	T	0.29	-0.295	0.04	2.1x10 ⁻¹¹	-0.147	0.06	7.7x10 ⁻³	244,143	-0.238	0.03	5.1x10 ⁻¹²		ADAMTS7 ^{cdefg}
chr15mb95	15	95,312,071	rs12906962	T	0.68	-0.292	0.04	5.3x10 ⁻¹²	-0.155	0.04	1.5x10 ⁻⁴	319,952	-0.221	0.03	5.6x10 ⁻¹⁴	SBP	LOC440311 ^{ef} , MCTP2 ^g
PPL	16	4,943,019	rs12921187	T	0.43	-0.203	0.04	3.0x10 ⁻⁷	-0.147	0.04	1.2x10 ⁻⁴	326,469	-0.174	0.03	2.5x10 ⁻¹⁰	SBP	PPL ^{aefg}
FBXL19	16	30,936,743	rs72799341	A	0.24	0.235	0.05	3.0x10 ⁻⁷	0.139	0.04	1.6x10 ⁻³	324,502	0.185	0.03	5.8x10 ⁻⁹		CTF1 ^{cdf} , FBXL19 ^{efg}
CMIP	16	81,574,197	rs8059962	T	0.42	-0.241	0.04	2.0x10 ⁻⁹	-0.103	0.04	8.5x10 ⁻³	319,839	-0.170	0.03	1.3x10 ⁻⁹		CMIP ^g
ACE	17	61,559,625	rs4308	A	0.37	0.242	0.04	3.2x10 ⁻⁹	0.186	0.04	2.7x10 ⁻⁶	319,394	0.213	0.03	6.8x10 ⁻¹⁴	SBP	ACE ^{cdefg}
MAPK4	18	48,142,854	rs745821	T	0.76	0.236	0.05	3.2x10 ⁻⁷	0.150	0.04	4.2x10 ⁻⁴	330,954	0.189	0.03	1.4x10 ⁻⁹		MAPK4 ^{defg}
CCNE1	19	30,294,991	rs62104477	T	0.33	0.209	0.04	7.1x10 ⁻⁷	0.148	0.04	2.4x10 ⁻⁴	320,347	0.177	0.03	1.2x10 ⁻⁹		CCNE1 ^{efg}
PLCB1	20	8,626,271	rs6108168	A	0.25	-0.305	0.05	1.5x10 ⁻¹¹	-0.127	0.04	2.9x10 ⁻³	327,368	-0.211	0.03	1.1x10 ⁻¹¹	SBP	PLCB1 ^{defg}
Pulse pressure																	
chr1mb9	1	9,441,949	rs9662255	A	0.43	-0.303	0.05	4.7x10 ⁻¹⁰	-0.130	0.04	3.0x10 ⁻³	310,618	-0.207	0.03	1.9x10 ⁻¹⁰		SPSB1 ^{efg}
SF3A3	1	38,455,891	rs4360494	C	0.55	0.332	0.05	5.7x10 ⁻¹²	0.224	0.05	3.6x10 ⁻⁶	282,851	0.278	0.03	3.7x10 ⁻¹⁶		SF3A3 ^{bfg} , FHL3 ^{bef}
RP4-710M16.1-PPAP2B	1	56,576,924	rs112557609	A	0.35	0.280	0.05	3.2x10 ⁻⁸	0.187	0.04	1.8x10 ⁻⁵	325,952	0.227	0.03	6.8x10 ⁻¹²	SBP	PLPP3 ^{cefg}
FGGY	1	59,653,742	rs3889199	A	0.71	0.462	0.05	3.3x10 ⁻¹⁸	0.271	0.05	1.9x10 ⁻⁹	329,486	0.351	0.03	1.8x10 ⁻²⁴	SBP	FGGY ^{dfig} , HSD52 ^{ef}
C2orf43	2	20,881,840	rs2289081	C	0.36	-0.251	0.05	5.3x10 ⁻⁷	-0.203	0.04	1.7x10 ⁻⁶	329,140	-0.223	0.03	5.5x10 ⁻¹²		GDF7 ^{efg}
PRKCE	2	46,363,336	rs11690961	A	0.88	0.437	0.07	4.2x10 ⁻⁹	0.266	0.07	4.6x10 ⁻⁵	327,847	0.340	0.05	3.9x10 ⁻¹²		PRKCE ^{dfig}
CEP68	2	65,283,972	rs74181299	T	0.62	0.296	0.05	2.1x10 ⁻⁹	0.181	0.04	2.0x10 ⁻⁵	324,224	0.230	0.03	9.6x10 ⁻¹³	SBP	CEP68 ^{efg}
TCF7L1	2	85,491,365	rs11689667	T	0.54	0.256	0.05	1.1x10 ⁻⁷	0.118	0.04	3.8x10 ⁻³	330,634	0.176	0.03	1.7x10 ⁻⁸		TCF7L1 ^{cefg}
FN1	2	216,300,482	rs1250259	A	0.74	-0.457	0.05	5.5x10 ⁻¹⁷	-0.210	0.05	7.7x10 ⁻⁶	325,485	-0.314	0.04	8.7x10 ⁻¹⁹	SBP	FN1 ^{cedfg}
GATA2	3	128,201,889	rs62270945	T	0.03	0.861	0.14	2.6x10 ⁻⁹	0.366	0.14	9.5x10 ⁻³	279,925	0.607	0.10	1.8x10 ⁻⁹		GATA2 ^{cefg}
PALLD	4	169,717,148	rs1566497	A	0.42	0.320	0.05	6.6x10 ⁻¹¹	0.173	0.04	4.8x10 ⁻⁵	320,948	0.236	0.03	1.9x10 ⁻¹³		PALLD ^{cdfig}
chr4mb174	4	174,584,663	rs17059668	C	0.92	-0.442	0.09	9.0x10 ⁻⁷	-0.245	0.08	2.2x10 ⁻³	313,277	-0.332	0.06	2.8x10 ⁻⁸		HAND2-AS1 ^g
LHFPL2	5	77,837,789	rs10057188	A	0.46	-0.280	0.05	5.5x10 ⁻⁹	-0.149	0.04	3.3x10 ⁻⁴	325,985	-0.205	0.03	6.7x10 ⁻¹¹	SBP	LHFPL2 ^{efg}
GJA1	6	121,781,390	rs11154027	T	0.47	0.311	0.05	1.1x10 ⁻¹⁰	0.125	0.04	3.7x10 ⁻³	316,708	0.207	0.03	1.1x10 ⁻¹⁰		GJA1 ^{cdfig}
ESR1	6	152,397,912	rs36083386	I	0.11	0.651	0.08	4.6x10 ⁻¹⁷	0.289	0.07	1.0x10 ⁻⁵	323,303	0.439	0.05	1.5x10 ⁻¹⁸		ESR1 ^{ecdfig}
FNDC1	6	159,699,125	rs449789	C	0.14	0.480	0.07	2.2x10 ⁻¹²	0.264	0.06	1.3x10 ⁻⁵	325,584	0.359	0.05	2.4x10 ⁻¹⁵		FNDC1 ^{defg}

THBS2	6	169,587,103	rs1322639	A	0.78	0.433	0.06	7.7x10 ⁻¹⁴	0.230	0.05	3.4x10 ⁻⁶	319,866	0.316	0.04	4.8x10 ⁻¹⁷		THBS2 ^{cdefg}
SUGCT	7	40,447,971	rs76206723	A	0.10	-0.405	0.08	2.6x10 ⁻⁷	-0.305	0.07	3.8x10 ⁻⁶	328,162	-0.346	0.05	7.4x10 ⁻¹²		SUGCT ^g
SLC20A2	8	42,324,765	rs2978456	T	0.55	-0.253	0.05	1.3x10 ⁻⁷	-0.130	0.05	4.4x10 ⁻³	304,964	-0.188	0.03	1.2x10 ⁻⁸		SLC20A2 ^{defg}
TRAPPC9	8	141,060,027	rs4454254	A	0.63	-0.320	0.05	9.4x10 ⁻¹¹	-0.217	0.04	2.9x10 ⁻⁷	330,022	-0.261	0.03	5.1x10 ⁻¹⁶		TRAPPC9 ^{efg}
SCAI	9	127,900,996	rs72765298	T	0.87	-0.392	0.07	5.9x10 ⁻⁸	-0.358	0.07	8.6x10 ⁻⁸	316,271	-0.374	0.05	2.7x10 ⁻¹⁴	SBP	RABEPK ^{af} , SCAI ^{efg}
KIAA1462	10	30,317,073	rs9337951	A	0.34	0.301	0.05	7.6x10 ⁻⁹	0.262	0.05	5.5x10 ⁻⁸	299,646	0.280	0.04	2.5x10 ⁻¹⁵		KIAA1462 ^{defg}
ARHGAP12	10	32,082,658	rs10826995	T	0.71	-0.317	0.05	2.2x10 ⁻⁹	-0.133	0.05	3.9x10 ⁻³	327,373	-0.212	0.03	1.1x10 ⁻⁹		ZEB1 ^{cf} , ARHGAP12 ^{efg}
PRDM11	11	45,208,141	rs11442819	I	0.11	-0.412	0.07	3.8x10 ⁻⁸	-0.185	0.06	3.3x10 ⁻³	326,483	-0.279	0.05	7.1x10 ⁻⁹		PRDM11 ^{efg}
NOX4	11	89,224,453	rs2289125	A	0.21	-0.481	0.06	3.1x10 ⁻¹⁶	-0.293	0.05	2.9x10 ⁻⁸	307,682	-0.377	0.04	9.1x10 ⁻²²		NOX4 ^{acdefg}
CEP164	11	117,283,676	rs8258	T	0.38	0.341	0.05	5.3x10 ⁻¹²	0.157	0.04	2.4x10 ⁻⁴	327,038	0.236	0.03	2.9x10 ⁻¹³		CEP164 ^{defg}
CCDC41	12	94,880,742	rs139236208	A	0.10	-0.442	0.08	5.7x10 ⁻⁸	-0.288	0.08	2.8x10 ⁻⁴	291,244	-0.363	0.06	1.6x10 ⁻¹⁰		CEP83-AS1 ^{ef} , CEP83 ^g
RP11-6101.1	14	98,587,630	rs9323988	T	0.63	-0.291	0.05	5.6x10 ⁻⁹	-0.156	0.04	2.0x10 ⁻⁴	327,551	-0.212	0.03	4.1x10 ⁻¹¹		C14orf177 ^g
VAC14	16	70,755,610	rs117006983	A	0.01	1.448	0.30	9.4x10 ⁻⁷	0.847	0.16	1.8x10 ⁻⁷	250,766	0.986	0.14	4.1x10 ⁻¹²		VAC14 ^{efg}
CDH13	16	83,045,790	rs7500448	A	0.75	0.386	0.06	4.2x10 ⁻¹²	0.288	0.05	1.8x10 ⁻⁹	321,958	0.329	0.04	1.1x10 ⁻¹⁹		CDH13 ^{bdefg}
KIAA0753	17	6,473,828	rs7226020	T	0.56	-0.348	0.05	1.3x10 ⁻¹²	-0.175	0.05	1.4x10 ⁻⁴	303,389	-0.256	0.03	2.3x10 ⁻¹⁴		KIAA0753 ^{bfg} , PITPNM3 ^{ef}
TP53-SLC2A4	17	7,571,752	rs78378222	T	0.99	1.530	0.22	8.9x10 ⁻¹²	0.487	0.18	7.9x10 ⁻³	294,053	0.904	0.14	1.8x10 ⁻¹⁰	DBP	TP53 ^{cdefg}
KCNH4-HSD17B1	17	40,317,241	rs79089478	T	0.97	0.842	0.15	1.2x10 ⁻⁸	0.377	0.13	4.4x10 ⁻³	318,326	0.584	0.10	3.1x10 ⁻⁹		KCNH4 ^g
PYY	17	42,060,631	rs62080325	A	0.66	-0.260	0.05	3.6x10 ⁻⁷	-0.128	0.05	4.8x10 ⁻³	315,689	-0.186	0.03	4.0x10 ⁻⁸		PYY ^{cefg}
MRC2	17	60,767,151	rs740698	T	0.56	-0.307	0.05	2.1x10 ⁻¹⁰	-0.161	0.04	2.8x10 ⁻⁴	311,450	-0.228	0.03	3.1x10 ⁻¹²		MRC2 ^{efg}
SLC14A2	18	43,097,750	rs7236548	A	0.18	0.462	0.06	1.1x10 ⁻¹³	0.273	0.05	2.2x10 ⁻⁷	330,075	0.352	0.04	2.0x10 ⁻¹⁸		SLC14A2 ^{cdefg}
SLC24A3	20	19,465,907	rs6081613	A	0.28	0.326	0.05	1.2x10 ⁻⁹	0.213	0.05	8.1x10 ⁻⁶	315,546	0.263	0.04	1.6x10 ⁻¹³		SLC24A3 ^{efg}
ARVCF	22	19,967,980	rs12628032	T	0.30	0.269	0.05	2.4x10 ⁻⁷	0.216	0.05	3.8x10 ⁻⁶	310,292	0.240	0.03	5.5x10 ⁻¹²	SBP	ARVCF ^{efg}
XRCC6	22	42,038,786	rs73161324	T	0.05	0.611	0.11	6.5x10 ⁻⁹	0.380	0.11	3.1x10 ⁻⁴	267,722	0.496	0.07	2.8x10 ⁻¹¹		XRCC6 ^g
(b) UK Biobank exome																	
Systolic blood pressure																	
SSPN	12	26,438,189	rs6487543	A	0.77	0.345	0.09	5.9x10 ⁻⁵	0.279	0.06	2.1x10 ⁻⁶	244,842	0.300	0.05	6.3x10 ⁻¹⁰	DBP	SSPN ^{dfg}
Diastolic blood pressure																	
MRAS	3	138,119,952	rs2306374	T	0.84	-0.237	0.05	9.3x10 ⁻⁶	-0.155	0.04	9.3x10 ⁻⁵	281,715	-0.184	0.03	7.4x10 ⁻⁹	SBP	MRAS ^{defg}
Pulse pressure																	

CD34	1	208,024,820	rs12731740	T	0.10	-0.360	0.08	5.8x10 ⁻⁶	-0.202	0.05	1.1x10 ⁻⁴	279,078	-0.249	0.04	1.1x10 ⁻⁸		MIR29B2 ^{df} , CD34 ^{dfg} , LOC148696 ^{ef}
ZNF638	2	71,627,539	rs3771371	T	0.57	-0.223	0.05	4.1x10 ⁻⁶	-0.130	0.03	9.6x10 ⁻⁵	280,285	-0.160	0.03	5.8x10 ⁻⁹		DYSF ^{df} , ZNF638 ^{efg}
CRACR2B	11	828,916	rs7126805	A	0.73	0.262	0.05	1.1x10 ⁻⁶	0.184	0.05	4.6x10 ⁻⁴	145,162	0.222	0.04	3.3x10 ⁻⁹		CD151 ^{cdfg} , CRACR2B ^{ef}

768

769 Locus: named according to the nearest annotated gene(s); Pos: build 37; EA: effect allele; EAF: effect allele frequency from discovery data in UK Biobank; Beta: effect
770 estimate from linear regression; SE: Standard Error of effect estimate; *P*: *P*-value of association; N: total sample size analysed; Traits: the other BP traits which reached
771 genome-wide significance in the combined meta-analysis. Note: within the UK Biobank discovery analysis sample size was N=140,882/140,886 for systolic and pulse
772 pressure / diastolic pressure, imputation quality score from SNPTEST ≥ 0.93 for all loci.

773 Candidate genes have been identified by one or multiple strategies:

774 ^acoding, nonsynonymous variant;

775 ^bGTEX eQTL

776 ^cCV KO Phenotype

777 ^dsupporting biology

778 ^eHi-C support

779 ^fvascular expression

780 ^gnearest to lead SNP

781 **Table 2: Association results for new independent secondary variants identified at (a) validated novel loci and (b) previously reported blood**
782 **pressure loci from either UK Biobank-GWAS or exome discovery.** All listed secondary variants were validated in the replication meta-analyses
783 and passed the conditional test for independence from the (a) sentinel novel variant from Table 1, or (b) previously reported SNVs (see
784 Supplementary Tables 8 and 10).

785

Secondary SNV in the locus						UK Biobank discovery					Replication			Combined			
Locus	Chr	Pos	rsID	EA	Trait	INFO	EAF	Beta	SE	P	Beta	SE	P	N	Beta	SE	P
(a) Validated novel loci from UK Biobank GWAS																	
<i>NADK-CPSF3L</i>	1	1254436	rs1886773	A	PP	0.99	0.03	-0.743	0.13	2.0x10 ⁻⁸	-0.481	0.15	1.0x10 ⁻³	233,789	-0.625	0.10	1.9x10 ⁻¹⁰
<i>RP4-710M16.1-PPAP2B</i>	1	56938218	rs6588634	T	PP	0.99	0.89	0.403	0.08	2.1x10 ⁻⁷	0.270	0.07	4.7x10 ⁻⁵	329,029	0.326	0.05	1.0x10 ⁻¹⁰
<i>FN1</i>	2	216245694	rs34923683	A	PP	1.00	0.02	0.837	0.15	4.8x10 ⁻⁸	0.432	0.16	7.7x10 ⁻³	285,653	0.646	0.11	6.8x10 ⁻⁹
<i>TP53-SLC2A4</i>	17	7185062	rs5417	A	DBP	0.99	0.57	0.207	0.04	2.1x10 ⁻⁷	0.207	0.04	1.1x10 ⁻⁷	319,299	0.207	0.03	1.1x10 ⁻¹³
<i>KCNH4-HSD17B1</i>	17	40709867	rs138643143	A	PP	0.85	0.07	0.539	0.10	1.4x10 ⁻⁷	0.420	0.15	5.8x10 ⁻³	229,161	0.502	0.08	3.3x10 ⁻⁹
(b) Previously reported loci																	
UK Biobank GWAS																	
<i>RNF207</i>	1	6683240	rs14057	A	SBP	0.99	0.35	-0.394	0.07	7.5x10 ⁻⁸	-0.235	0.06	2.0x10 ⁻⁴	329,584	-0.303	0.05	2.5x10 ⁻¹⁰
<i>FIGN-GRB14</i>	2	165513065	rs34271465*	D	SBP	1.00	0.41	-0.370	0.07	1.9x10 ⁻⁷	-0.277	0.06	6.9x10 ⁻⁶	328,486	-0.317	0.05	9.9x10 ⁻¹²
<i>ENPEP</i>	4	111431444	rs33966350*	A	SBP	1.00	0.01	1.742	0.31	2.6x10 ⁻⁸	1.525	0.41	1.8x10 ⁻⁴	216,630	1.661	0.25	2.1x10 ⁻¹¹
<i>GUCY1A3-GUCY1B3</i>	4	156406054	rs146853253*	D	PP	0.99	0.16	0.457	0.06	1.7x10 ⁻¹²	0.212	0.06	1.4x10 ⁻⁴	322,302	0.316	0.04	6.9x10 ⁻¹⁴
<i>EBF1</i>	5	158220193	rs31864	A	PP	0.99	0.55	0.307	0.05	1.9x10 ⁻¹⁰	0.132	0.04	1.5x10 ⁻³	326,557	0.206	0.03	5.5x10 ⁻¹¹
<i>EBF1</i>	5	158448401	rs888987	C	DBP	0.96	0.37	0.208	0.04	4.4x10 ⁻⁷	0.111	0.04	7.1x10 ⁻³	311,814	0.160	0.03	4.3x10 ⁻⁸
<i>PDE3A</i>	12	19979881	rs10841376	C	SBP	0.99	0.76	0.261	0.08	1.6x10 ⁻³	0.362	0.07	5.1x10 ⁻⁷	327,370	0.319	0.05	4.5x10 ⁻⁹
<i>PDE3A</i>	12	20230639	rs10770612	A	PP	1.00	0.80	0.378	0.06	2.5x10 ⁻¹⁰	0.259	0.05	1.8x10 ⁻⁶	311,586	0.313	0.04	6.9x10 ⁻¹⁵
<i>PDE3A</i>	12	20368269	rs60691990*	T	DBP	0.98	0.65	0.344	0.04	1.4x10 ⁻¹⁶	0.223	0.04	7.4x10 ⁻⁸	323,722	0.283	0.03	5.0x10 ⁻²²
<i>TBX5-TBX3</i>	12	115928440	rs10850519*	C	DBP	0.99	0.30	-0.244	0.04	1.4x10 ⁻⁸	-0.188	0.04	4.7x10 ⁻⁶	327,837	-0.214	0.03	5.1x10 ⁻¹³
<i>MYH6</i>	14	23761094	rs12050260	T	PP	0.97	0.35	0.261	0.05	2.9x10 ⁻⁷	0.132	0.05	4.1x10 ⁻³	304,390	0.190	0.03	2.6x10 ⁻⁸
<i>FURIN-FES</i>	15	91427692	rs138682554	A	SBP	0.85	0.03	1.274	0.23	5.1x10 ⁻⁸	0.695	0.21	8.8x10 ⁻⁴	279,876	0.952	0.16	9.8x10 ⁻¹⁰
<i>HOXB7</i>	17	46874272	rs585736	A	PP	1.00	0.03	0.712	0.13	7.8x10 ⁻⁸	0.517	0.13	4.1x10 ⁻⁵	301,845	0.609	0.09	2.5x10 ⁻¹¹
<i>INSR</i>	19	7258405	rs11671314	C	SBP	0.94	0.13	0.532	0.11	8.3x10 ⁻⁷	0.344	0.13	6.2x10 ⁻³	253,103	0.452	0.08	3.4x10 ⁻⁸
<i>JAG1</i>	20	10669188	rs2206815	A	PP	0.98	0.50	-0.432	0.05	3.9x10 ⁻¹⁹	-0.247	0.04	2.7x10 ⁻⁹	324,088	-0.326	0.03	4.7x10 ⁻²⁵
<i>JAG1</i>	20	10767811	rs1040922	T	DBP	0.99	0.28	-0.344	0.04	3.8x10 ⁻¹⁵	-0.156	0.04	1.8x10 ⁻⁴	325,879	-0.245	0.03	4.2x10 ⁻¹⁶

<i>PREX1</i>	20	47411149	rs80346118	A	DBP	0.99	0.15	-0.305	0.06	3.1x10 ⁻⁸	-0.243	0.05	5.6x10 ⁻⁶	327,614	-0.273	0.04	1.1x10 ⁻¹²
<i>CRYAA-SIK1</i>	21	44720890	rs79094191	T	DBP	0.98	0.96	-0.691	0.10	3.9x10 ⁻¹¹	-0.408	0.12	4.4x10 ⁻⁴	284,734	-0.564	0.08	3.8x10 ⁻¹³
UK Biobank exome																	
<i>ST7L-CAPZA1-MOV10</i>	1	113456546	rs1049434*	A	DBP	1.00	0.44	-0.175	0.04	9.7x10 ⁻⁶	-0.131	0.03	1.1x10 ⁻⁵	264,717	-0.147	0.02	6.6x10 ⁻¹⁰
<i>CDH17</i>	8	95264265	rs138582164	A	PP	0.78	0.001	5.199	0.99	1.3x10 ⁻⁷	2.620	0.73	3.2x10 ⁻⁴	226,592	3.529	0.59	1.7x10 ⁻⁹

786

787 Locus: For (a) the locus name from Table 1 for the nearest annotated gene, (b) the name of the previously reported blood pressure locus; Pos: build 37; EA: effect allele;
788 Trait: the validated trait with most significant association in the combined meta-analysis; INFO: imputation quality score; EAF: effect allele frequency from discovery data in
789 UK Biobank; Beta: effect estimate from linear regression; SE: Standard Error of effect estimate; *P*: *P*-value of association; N: total sample size analysed; (Note: within the UK
790 Biobank discovery analysis the sample size was N=140,882/140,886 for systolic and pulse pressure / diastolic pressure.) The variants with * denotes secondary signals
791 which are in LD ($r^2 \geq 0.2$) with secondary signals which have been published since the time of our study^{10,11}

Figure 1: Study design schematic for discovery and validation of novel loci. N: sample size; QC: Quality Control; PCA: Principal Component Analysis; BP: blood pressure; SBP: systolic BP; DBP: diastolic BP; PP: pulse pressure; SNVs: single nucleotide variants; BMI: body mass index; UKB: UK Biobank; UKBL: UK BiLEVE; GWAS: Genome-wide association study; MAF: Minor Allele Frequency; *P*: P-value; LD: Linkage Disequilibrium; 1000G: 1000 Genomes. UKBBvsUKBL: a binary indicator variable for UK Biobank vs UK BiLEVE to adjust for the different genotyping chips

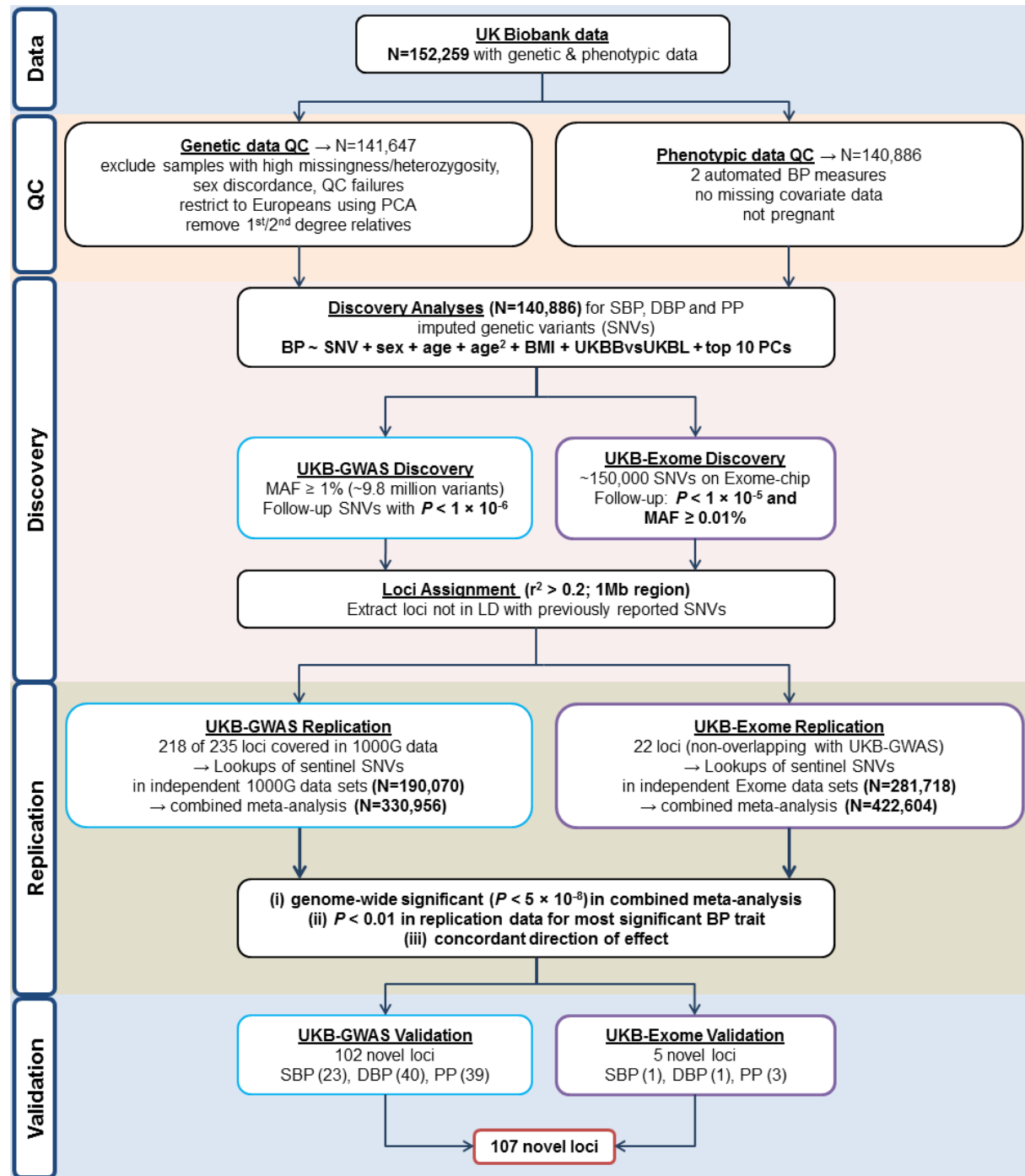


Figure 2: UK Biobank GWAS discovery Venn diagram of 107 validated novel loci showing concordance of significant associations across the three blood pressure phenotypes for the 107 novel sentinel variants (Table 1) from both the GWAS and exome analyses, according to genome-wide significance in the combined meta-analysis. The locus names labelled within the Venn Diagram correspond to Table 1, and relate to the nearest annotated gene.

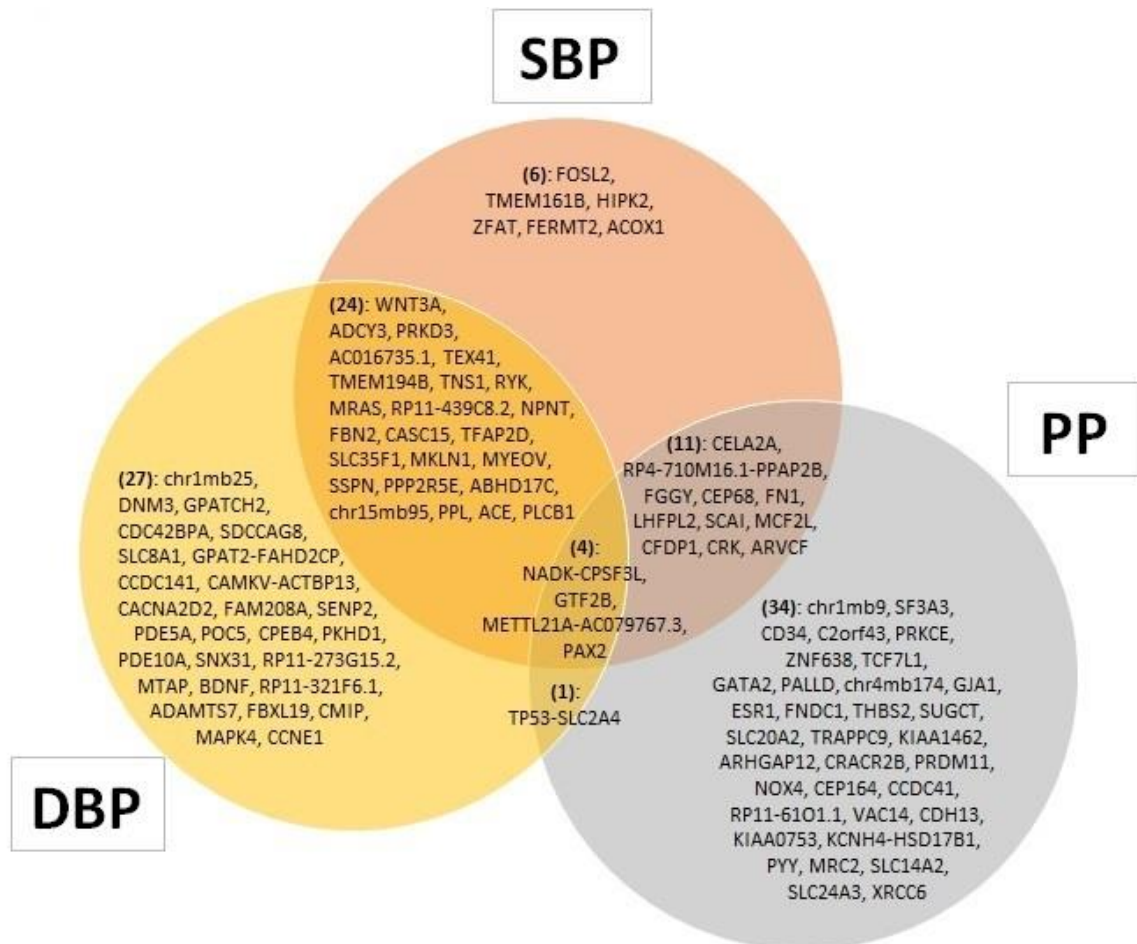


Figure 3: Association of blood pressure loci with other traits. Plot (A) shows results for associations with other traits which were extracted from the PhenoScanner database for the sentinel novel variants from Table 1, including proxies in Linkage Disequilibrium ($r^2 \geq 0.8$), with genome-wide significant associations ($P < 5 \times 10^{-8}$). The loci are grouped by blood pressure traits ordered right to left according to the loci in Table 1. There are four systolic blood pressure associated loci, 14 diastolic blood pressure associated loci and nine pulse pressure associated loci with associations with other traits reported in the literature. Traits are grouped into different disease categories: “Pulse/HR” includes pulse, heart rate, pulse wave velocity and aortic stiffness traits; “CAD/CHD/MI”: Coronary Artery Disease / Coronary Heart Disease / Myocardial Infarction; “Blood” traits: Haemoglobin levels and platelet counts; “Lipids”: LDL

and Total Cholesterol; “BMI/WHR” includes Body Mass Index, weight, obesity, waist or hip circumference, Waist-Hip-Ratio; “Menarche”: age at menarche; “Lung”: lung function (FEV1); “Alzheimer’s” traits refers to Cerebrospinal fluid levels of Alzheimer’s disease related proteins; “Cancer” includes carcinomas, neuroblastomas, bladder cancer; “Education”: years of educational attainment.

Plots (B) and (C) show mouse phenotype enrichment and disease ontology enrichment, respectively, of validated novel and previously reported variants. Enrichment was performed using the GREAT tool (<http://bejerano.stanford.edu/great>) with the sentinel SNVs as query.

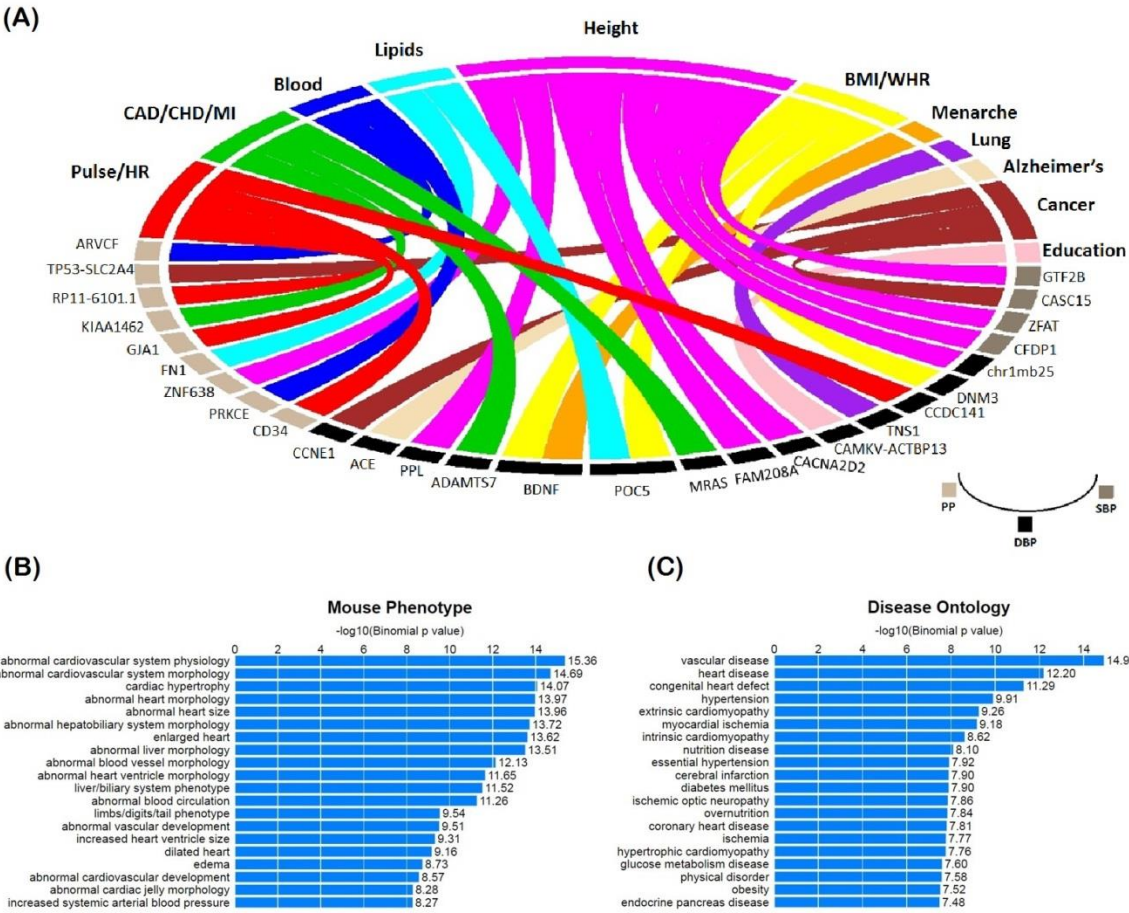


Figure 4: Distribution of Genetic Risk Score (GRS) based on previously reported and validated novel blood pressure variants and its relationship with blood pressure values, hypertension and CVD outcomes. A, Distribution of GRS in Airwave and sex-adjusted odds ratio of hypertension in age 50+ comparing each of the upper four GRS quintiles with the lowest quintile; dotted lines represent the upper 95% confidence intervals. B, Mean blood pressures and standard deviation in bracket in Airwave age 50+ across GRS quintiles. C, Distribution of GRS in UKB and sex-adjusted odds ratio of CVD, CAD and stroke comparing each of the upper four GRS quintiles with the lowest quintile; dotted lines represent the upper 95% confidence intervals. D, Count of CVD, CAD and stroke (events and deaths) across GRS quintiles in UKB participants

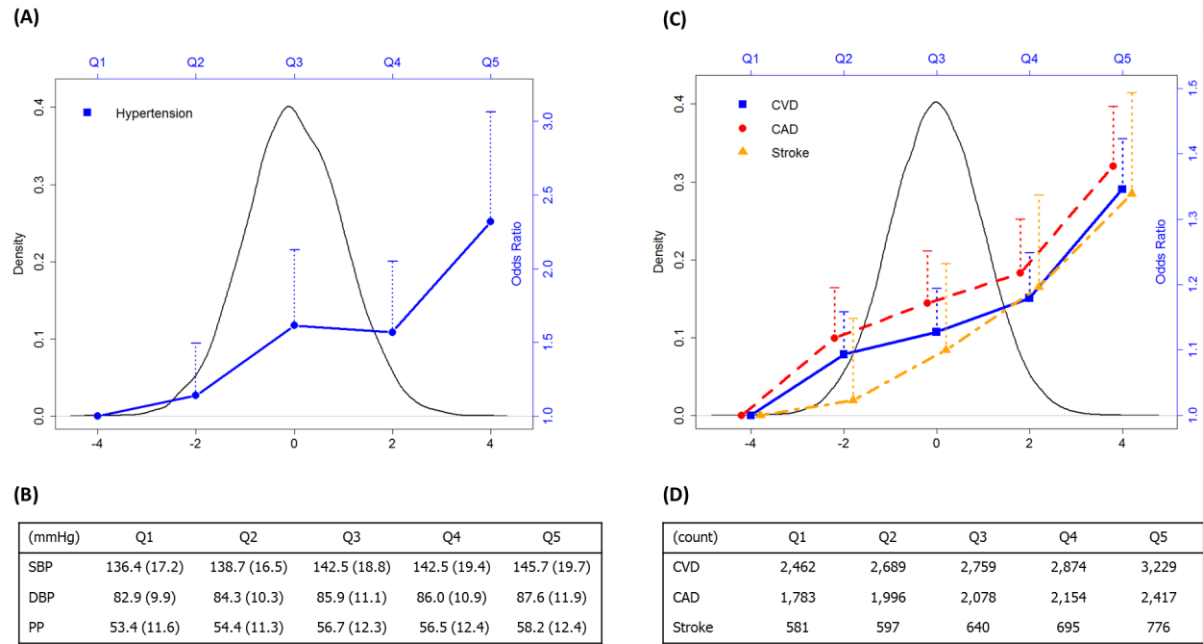
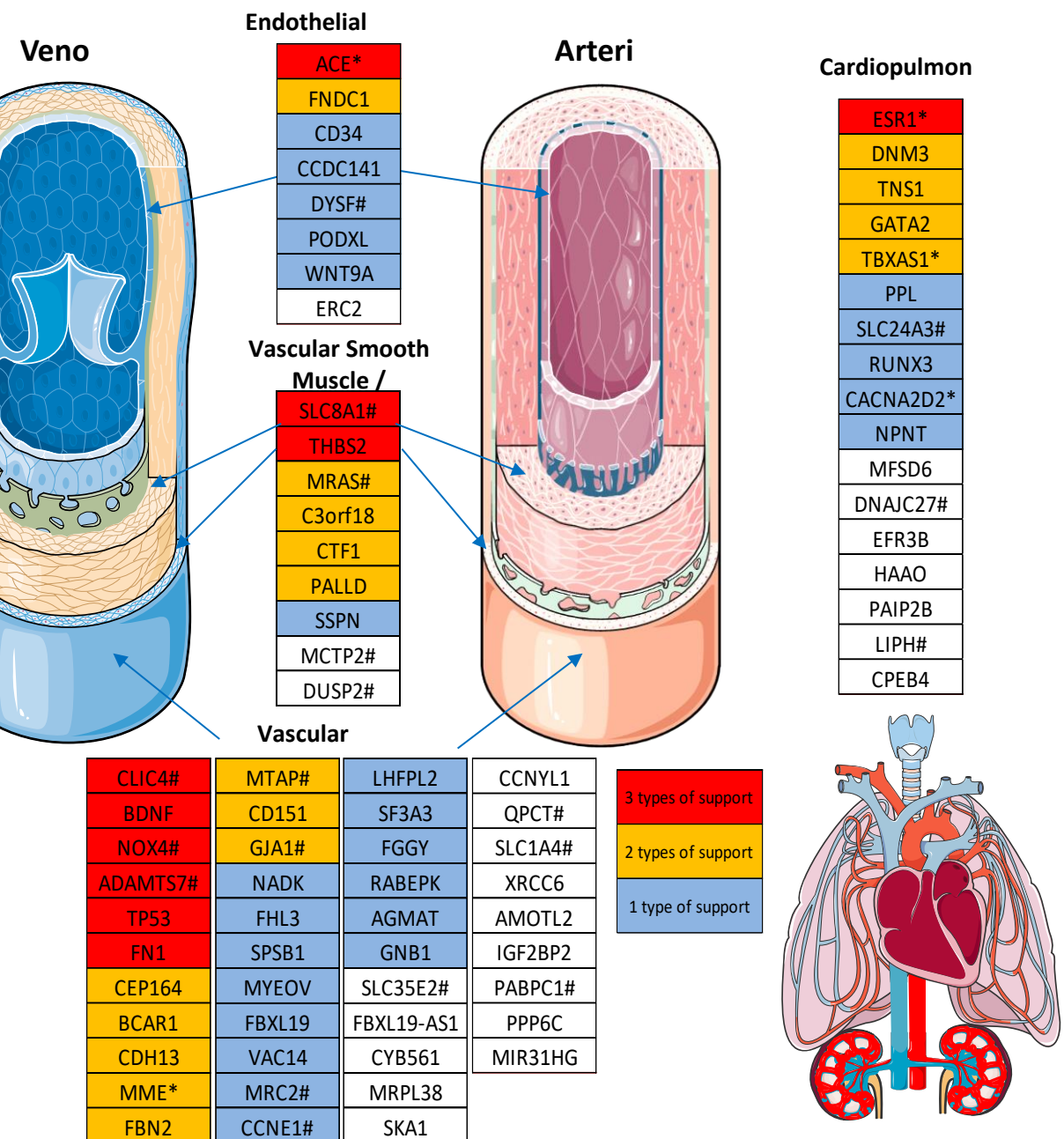


Figure 5: Summary of novel gene cardiovascular expression. Genes are shown on the basis of their tissue expression and supporting evidence summarised in Supplementary Table 14, based on Knockout (KO) phenotype, previously reported blood pressure biology or a strong functional rationale: eQTL (expression Quantitative Trait Loci), nsSNV (non-synonymous SNV), Hi-C. Multiple lines of evidence indicate the central importance of the vasculature in blood pressure regulation and we thus highlight existing drugged (*) and druggable (#) targets among these genes. Illustrations used elements with permission from Servier Medical Art: www.servier.fr/servier-medical-art. We note that some druggable genes may carry a safety liability, such as GJA1, which has known association with QT interval²³



Online Methods

UK Biobank data

Our Genome Wide Association Study (GWAS) analysis is performed using data from the interim release of the first ~150k UK Biobank participants (Supplementary Methods)¹⁷. These consist of ~100k individuals from UK Biobank genotyped at ~800,000 single nucleotide variants (SNVs) with a custom Affymetrix UK Biobank Axiom Array chip⁶⁸ and ~50k individuals genotyped with a custom Affymetrix UK BiLEVE Axiom Array chip from the UK BiLEVE study⁶⁹, which is a subset of UK Biobank. SNVs were imputed centrally by UK Biobank using a merged UK10K sequencing + 1000 Genomes imputation reference panel.

Quality control

Following quality control (QC) procedures already carried out centrally by UK Biobank, we exclude discordant SNVs and samples with QC failures, gender discordance and high heterozygosity/missingness. We further restrict our data to a subset of individuals of European ancestry. By applying *kmeans* clustering to the Principal Component Analysis (PCA) data a total of N=145,315 Europeans remain. Then we use the kinship data to exclude 1st and 2nd degree relatives, with N=141,647 unrelated individuals remaining. Finally we restrict our data to non-pregnant individuals with two automated BP measurements available, resulting in a maximum of N=140,886 individuals for analysis (Supplementary Methods).

Phenotypic data

After calculating the mean systolic and diastolic pressure values from the two blood pressure measurements, we adjust for medication use by adding 15 and 10 mmHg to systolic and diastolic pressure, respectively, for individuals reported to be taking blood pressure-lowering medication (21.4% of individuals)⁷⁰. Pulse Pressure is calculated as systolic minus diastolic pressure, according to the medication-adjusted traits. Hypertension, used in secondary analyses, is defined as: (i) systolic pressure ≥ 140 mmHg, or (ii) diastolic pressure ≥ 90 mmHg, (iii) or taking blood pressure-lowering medication; otherwise individuals are classified as non-hypertensive. Descriptive summary statistics are provided for all individuals, and stratified by UK Biobank vs UK BiLEVE participants (**Supplementary Table 1**).

Analysis models

For the GWAS, we perform linear regression analyses of the three (untransformed) continuous, medication-adjusted BP traits (systolic, diastolic and pulse pressure) for all measured and imputed genetic variants in dosage format using SNPTEST software⁷¹ under an additive genetic model. We carry out a similar analysis for the exome content. Each analysis includes the following covariates: sex, age, age², body mass index, top ten PCs and a binary indicator variable for UK Biobank vs UK BiLEVE to adjust for the different genotyping chips. We also run an association analysis within UK Biobank for validated novel blood pressure SNVs and hypertension using logistic regression under an additive model with adjustments as above. There are 76,554 hypertensive cases and the 64,384 remaining participants are

treated as non-hypertensive controls. This sample size is slightly larger than the N=140,866 used in the main analyses, since participants with only one blood pressure measurement, but with reported blood pressure-lowering medication, could be included as hypertensive.

Previously reported variants

We compile a list of all SNVs previously reported to be associated with blood pressure (**Supplementary Table 12**). This list includes all published SNVs which have been identified and validated from previous GWAS, CardioMetaboChip and exome chip projects¹⁰⁻¹². We augment this list to include all 34,459 SNVs in Linkage Disequilibrium (LD) with the previously reported SNVs, according to a threshold of $r^2 \geq 0.2$. Results for all these variants are extracted for each of the three blood pressure traits, to check previously reported blood pressure associations in the UK Biobank data, according to whether the sentinel SNV or a variant at the locus in LD ($r^2 \geq 0.2$) with it showed evidence of support ($P < 0.01$) for association with at least one of the three BP traits.

Replication strategy

We use three independent external data sets for replication (Supplementary Methods). First, for the GWAS analysis based on advanced 1000 Genomes imputation enhanced by UK10K data we consider SNVs with MAF $\geq 1\%$ and perform a reciprocal replication exchange with the International Consortium of Blood Pressure (ICBP) 1000 Genomes meta-analysis (max N = 150,134). The imputation strategy for ICBP 1000 Genomes meta-analysis is based on an earlier imputation grid for the 1000 Genomes project. In addition, we recruit further cohorts with 1000 Genomes data which had not contributed to the ICBP-1000 Genomes discovery meta-analysis: ASCOT-UK (N = 3,803), ASCOT-SC (N = 2,462), BRIGHT (N = 1,791), Generation Scotland (GS) (N = 9,749), EGCUT (N = 5,468), Lifelines (N = 13,292) and PREVEND (N = 3,619). This gives a total of N = 190,318 independent replication samples for the GWAS discovery.

Second, because the UK Biobank and UK BiLEVE genotyping chips contain exome content, we sought replication from two blood pressure exome consortia (European exome consortium and the Cohorts for Heart and Ageing research in Genome Epidemiology – CHARGE BP exome consortium), to allow validation of coding variants and variants with lower frequency. The European exome consortium (N = 161,926) and CHARGE consortium (N = 119,792) give a total of N = 281,718 independent replication samples for the UK Biobank exome discovery.

Note that the lookups for GWAS and exome discovery are distinct sets of SNVs. Loci are assigned sequentially, prioritising the primary GWAS discovery first, then considering any remaining loci with non-overlapping exome content for replication in the independent exome replication resources.

Statistical criteria for replication

For the GWAS discovery, there are ~9.8 million SNVs with MAF $\geq 1\%$ and INFO > 0.1 . We consider for follow-up any SNVs with $P < 1 \times 10^{-6}$ for any of the three blood pressure traits. For the exome discovery, there are 149,026 exome SNVs (Supplementary Methods) which were polymorphic with INFO > 0.1 ; for follow-up we consider all SNVs with MAF $\geq 0.01\%$ and $P < 1 \times 10^{-5}$. All such SNVs are annotated to loci according to both an LD threshold of $r^2 \geq 0.2$ and a

1Mb interval region (see Supplementary Methods), and signals are classified either as belonging to novel loci, or being potential secondary signals at previously reported loci.

Selection of variants for follow-up

The sentinel (most significant) SNV from each association signal is selected for follow-up, all of which are pairwise-independent by LD ($r^2 < 0.2$). For the GWAS discovery, we check that potential lookup SNVs are covered within the ICBP-1000G replication data (Supplementary Methods). Of the 235 novel loci containing previously unreported SNVs with $MAF \geq 1\%$, $INFO > 0.1$ and $P < 1 \times 10^{-6}$, 218 are covered, and similarly 100 of the 123 potential secondary SNVs at 51 of the 54 previously reported BP loci are available for follow-up. For the exome discovery, by following up SNVs with $MAF \geq 0.01\%$, $INFO > 0.1$ and $P < 1 \times 10^{-5}$ across the three blood pressure traits, we carry forward for replication sentinel SNVs at 22 novel loci, and potential secondary SNVs at three previously reported loci. We produce locus zoom plots for each of the lookup variants.

Replication meta-analyses

The replication and combined meta-analyses are performed within METAL software⁷² using fixed effects inverse variance weighted meta-analysis (Supplementary Methods). The combined meta-analysis of both the UK Biobank discovery ($N = 140,886$) and GWAS replication meta-analysis (max $N = 190,070$) include a total maximum sample size of $N = 330,956$. For the exome combined meta-analysis, we synthesize data from the UK Biobank discovery exome content (max $N=140,866$), with the replication dataset from both exome consortia (total max $N=281,718$), giving a maximum sample size of $N=422,604$.

Validation Criteria

In our study a signal is declared validated if it satisfies ALL of the following three criteria:

- (i) the sentinel SNV is genome-wide significant ($P < 5 \times 10^{-8}$) in the combined meta-analysis for any of the three blood pressure traits;
- (ii) the sentinel SNV shows evidence of support ($P < 0.01$) in the replication meta-analysis alone for association with the most significantly associated blood pressure trait from the combined meta-analysis;
- (iii) the sentinel SNV has concordant direction of effect between the UK Biobank discovery and the replication meta-analysis for the most significantly associated blood pressure trait from the combined meta-analysis.

Secondary signals

By conditional analysis within UK Biobank data we assess all validated secondary signals from novel and previously reported loci for independence from the sentinel or previously reported SNV, respectively (Supplementary Methods). We declare a secondary signal to be independent of the previously reported SNV if there is less than a 1.5 fold difference between the main association and conditional association P -values on a $-\log_{10}$ scale, i.e. if $-\log_{10}(P) / -\log_{10}(P_{\text{cond}}) < 1.5$. Note that the lookup criteria already ensure that the secondary variant

is not in LD ($r^2 < 0.2$) with the previously reported SNV. If more than one SNV in a region is found to be independent we undertake further rounds of iterative conditional analysis.

Lookups in non-European ancestries

As a secondary analysis, we look up 102 and 5 validated novel SNVs from the UK Biobank-GWAS and exome analyses, respectively, in non-European ancestry samples. These comprise analysis of East Asian (N = 31,513) and South Asian (N = 33,115) ancestry data from the iGEN-BP consortium¹³ for the GWAS lookups, and South Asian (N = 25,937), African American (N = 21,488) and Hispanic (N = 4,581) ancestry data from the CHARGE BP exome consortium¹² and CHD+ Exome consortium¹¹, for the exome content lookups (Supplementary Methods). We carry out a binomial (sign) test based on the number of SNVs with consistent directions of effect between UK Biobank and each of the non-European ancestry samples.

Monogenic blood pressure gene lookups

The UK Biobank and UK BiLEVE arrays include some rare coding variants for monogenic disorders. We collate a list of all specific mutation variants within genes known to be associated with monogenic blood pressure disorders²². Results from the UKB discovery association analyses for all three blood pressure traits are extracted for any of these SNVs directly covered within the UK Biobank dataset (**Supplementary Table 13**). Note that a search of proxies did not augment the list of available variants, so results are reported for the specific variants only.

Functional analyses

In order to prioritise associated SNVs, we use an integrative bioinformatics approach to collate functional annotation at both the variant and gene level for each SNV within the blood pressure loci (all SNVs in LD $r^2 \geq 0.8$ with the blood pressure-associated SNVs). At the variant level we use ANNOVAR⁷³ to obtain comprehensive functional characterisation of variants, including gene location, conservation and amino acid substitution impact based on a range of prediction tools including SIFT and polyphen2. All nonsynonymous variants were predicted damaging by two or more methods.

We use the University of California Santa Cruz (UCSC) genome browser to review sequence specific context of SNVs in relation to function, particularly in the Encyclopedia of DNA Elements (ENCODE) dataset⁷⁴. We use the UCSC table browser to annotate SNVs in ENCODE regulatory regions. We evaluate SNVs for impact on putative micro RNA target sites in the 3' un-translated regions (3'UTR) of transcripts by a query of the miRNASNP database⁷⁵. We evaluate all SNVs in LD ($r^2 \geq 0.8$) with our novel sentinel SNVs for evidence of mediation of expression quantitative trait loci (eQTL) in all 44 tissues using the Genotype-Tissue Expression (GTEx) database (www.gtexportal.org), in order to identify novel loci which are highly expressed, and to highlight specific tissue types which show eQTLs for a large proportion of novel loci. We further seek to identify novel loci with the strongest evidence of eQTL associations in arterial tissue, in particular.

At the gene level, we use Ingenuity Pathway Analysis (IPA) software (IPA®, QIAGEN Redwood City, www.qiagen.com/ingenuity) to review genes with prior links to blood pressure, based on

annotation with the “Blood Pressure” Medline Subject Heading (MESH) term which is annotated to 684 genes. We also use IPA to identify genes which interact with blood pressure MESH annotated genes, and evaluate genes for evidence of small molecule druggability based on queries of ChEMBL (www.ebi.ac.uk/chembl/) and Drug Gene Interaction database (dgidb.genome.wustl.edu).

We then perform overall enrichment testing across all loci. Firstly, we use DEPICT⁷⁶ (Data-driven Expression Prioritized Integration for Complex Traits) to identify highly expressed tissues and cells within the blood pressure loci. DEPICT uses a large number of microarrays (~37k) to identify cells and tissues where the genes are highly expressed and uses precomputed GWAS phenotypes to adjust for co-founding sources. DEPICT provides a *P*-value of enrichment and false discovery rates adjusted *P*-values for each tissue/cells tested.

Furthermore, to investigate regulatory regions, we employ a two tiered approach to investigate cell type specific enrichment within DNase I sites using FORGE, which tests for enrichment of SNVs within DNase I sites in 123 cell types from the Epigenomics Roadmap Project and ENCODE⁷⁷ (Supplementary Methods). Validated novel sentinel SNVs discovered in our study are analysed along with previously reported SNVs and secondary signals (with *P*-value < 1×10^{-4}) to evaluate the overall tissue specific enrichment of blood pressure associated variants. In a second analysis we use FORGE (with no LD filter) to investigate directly our curated candidate regulatory SNVs for overlap with cell-specific DNase I signals.

GenomeRunner⁷⁸ is used to search for enrichment of validated novel and previously reported sentinel SNVs with histone modification mark genomic features (Supplementary Methods). Relevant cardiovascular tissue expression is investigated using Fantom5 reference transcript expression data (fantom.gsc.riken.jp/5) (Supplementary Methods).

We use IPA (IPA®, QIAGEN Redwood City, www.qiagen.com/ingenuity) to identify biological pathways and transcriptional upstream regulators enriched for genes within the blood pressure loci. The transcriptional upstream regulator analysis aims to identify transcription factors, compounds, drugs, kinases and other molecules, for which the target is one of the blood pressure genes under investigation.

We query SNVs against PhenoScanner¹⁹ to investigate trait pleiotropy, extracting all association results with nominal significance at *P* < 0.05 for full reporting (**Supplementary Table 15**), and then extract genome-wide significant results to highlight the validated novel loci with strongest evidence of association with other traits (**Fig. 3a**). We also use the Genomic Regions Enrichment of Annotations Tool (GREAT) to study gene set enrichment of mouse phenotype and disease ontology terms within our validated novel and previously reported loci, using default SNV to gene mapping settings⁷⁹.

We carry out metabolomics analysis using two sets of data. First we use ¹H NMR lipidomics data on plasma from a subset of 2,000 participants of the Airwave Health Monitoring Study^{80,81} (Supplementary Methods). For each replicated blood pressure-associated SNV we ran association tests with the lipidomics data using linear regression analyses, adjusted for age and sex. We computed significance thresholds using a permutation derived family wise error rate (5%) to account for the high correlation structure of these data (ENT=35)⁸². We also

test each replicated SNV against published genome-wide vs metabolome-wide associations in plasma and urine using publicly available data from the “Metabolomics GWAS Server” to identify metabolites that have been associated with variants of interest at $P < 3.0 \times 10^{-4}$ (Bonferroni corrected P for validated signals)^{25,26}.

Experimental methods

We prioritise novel genes for laboratory testing on the basis of evidence for SNV function (including coding variants, eQTLs and Hi-C interactions), biological support for relevance to blood pressure (from literature review) and transgenic phenotype. We perform genotyping and Quantitative Reverse-Transcription Polymerase Chain Reaction (q RT-PCR) for the selected sentinel variants of interest using human vascular smooth muscle cells and endothelial cells and test for expression levels (Supplementary Methods). All three SNVs were tested using an additive model.

Genetic risk scores

First, by calculating genetic risk scores (GRS), we use the Airwave study⁸⁰ data to assess the effect in an independent cohort of the blood pressure-associated variants on blood pressure and risk of hypertension (Supplementary Methods). This provides an estimate of the combined effect of the blood pressure raising variants avoiding bias by “winners curse”. We create weighted GRSs for all pairwise-independent, LD-filtered ($r^2 < 0.2$) previously reported variants and validated novel variants (sentinel and secondary SNVs) combined, using SNVs available in Airwave (**Supplementary Table 21**). For the previously reported variants, we weight blood pressure increasing alleles by the beta coefficients from the UK Biobank discovery GWAS. For the novel variants, beta coefficients of the replication meta-analysis are used as independent, unbiased weights.

For the analyses of trait variance explained, we use three trait-specific GRSs (i.e. systolic, diastolic and pulse pressure). Each GRS includes all variants, but weights are trait-specific, using the beta coefficients from the analysis of each of the three different blood pressure traits, e.g. the systolic GRS is weighted by the beta coefficients from the systolic GRS. To calculate the percent of variance for each blood pressure trait explained by its corresponding trait-specific GRS, not accounted for by known factors, we generate the residuals from the regression model of each trait against covariates of age, age², sex and body mass index. We then fit a second linear model for the trait residuals with all the variants in the GRS plus the top 10 principal components. We calculate these percentage variance explained results within an independent population (Airwave).

For risk score analyses we calculate a single blood pressure GRS, as the average of the systolic and diastolic pressure GRSs. We standardize the average GRS to have mean of zero and standard deviation of one. We assess the association of the continuous average GRS variable with each blood pressure trait by simple linear regression. We also run a logistic regression to examine the association of the average GRS with risk of hypertension. We perform each analysis both with and without adjustment for sex. We test for interaction between age (below age 50, and 50 years and above) and the effect of the GRS on blood pressure. We then compare blood pressure levels and risk of hypertension for individuals in the top and bottom

20% of the GRS distribution at ages 50 years and over using linear and logistic regression, respectively.

We also assess the association of the average blood pressure GRS with cardiovascular outcomes in the UK Biobank data, based on self-reported medical history, and linkage to hospitalization and mortality data. We include all pairwise-independent previously reported blood pressure variants and validated novel variants. We use logistic regression with binary outcome variables for coronary heart disease, stroke and cardiovascular disease (see Supplementary Methods) and GRS as explanatory variable (with and without sex adjustment).

URLs

FORGE (accessed 16 Aug 2016),
http://browser.1000genomes.org/Homo_sapiens/UserData/Forge?db=core

Fantom5 data (accessed 16 Aug 2016), <http://fantom.gsc.riken.jp/5/>

ENCODE DNase I data (wgEncodeAwgDnaseMasterSites; accessed 20 Aug 2016 using Table browser)

ENCODE cell type data (accessed 20 Aug 2016),
<http://genome.ucsc.edu/ENCODE/cellTypes.html>.

Exome chip design:
http://genome.sph.umich.edu/wiki/Exome_Chip_Design

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